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I, KIM MARSHALL, MANAGER EXAMINATION SUPPORT AND SALES, hereby certify that the annexed is a true copy of the Provisional specification in connection with Application No. PP 0585 for a patent by COMMONWEALTH SCIENTIFIC AND INDUSTRIAL RESEARCH ORGANISATION filed on 27 November 1997.



WITNESS my hand this Ninth day of December 1998

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## **AUSTRALIA**

## Patents Act 1990

## COMMONWEALTH SCIENTIFIC AND INDUSTRIAL RESEARCH ORGANISATION

## PROVISIONAL SPECIFICATION

Invention Title:

Receptor agonists and antagonists

The invention is described in the following statement:

### **RECEPTOR AGONISTS AND ANTAGONISTS**

## Field of the Invention

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This invention relates to the field of receptor structure and receptor/ligand interactions. In particular it relates to the field of using receptor structure to predict the structure of related receptors and to use the determined structures and predicted structures to select and screen for agonists and antagonists of the polypeptide ligands.

## **Background of the Invention**

Insulin is the peptide hormone that regulates glucose uptake and metabolism. The two types of diabetes are associated with either an inability to produce insulin because of destruction of the pancreatic islet cells (Homo-Delarche, F. & Boitard, C.,1996, Immunol. Today 10: 456-460) or poor glucose metabolism resulting from either insulin resistance at the target tissues, inadequate insulin secretion by the islets or faulty liver function (Taylor, S. I., et al., 1994, Diabetes, 43: 735-740).

Insulin-like growth factors-1 and 2 (IGF-1 and 2) are structurally related to insulin but are more important in tissue growth and development than in metabolism. They are primarily produced in the liver in response to growth hormone but are also produced in most other tissues where they function as paracrine/autocrine regulators. The IGFs are strong mitogens and are involved in numerous physiological states and certain cancers (Baserga, R., 1996, TibTech 14: 150-152).

Epidermal growth factor (EGF) is a small polypeptide cytokine that is unrelated to the insulin/IGF family. It stimulates marked proliferation of epithelial tissues and is a member of a larger family of structurally related cytokines such as transforming growth factor α, amphiregulin, betacellulin, heparin-binding EGF and some viral gene products. Abnormal EGF family signalling is a characteristic of certain cancers (Soler, C. & Carpenter, G., 1994 In Nicola, N. (ed) Guidebook to Cytokines and Their receptors", Oxford Univ. Press, Oxford, pp194-197; Walker, F. & Burgess, A. W., 1994, In Nicola, N. (ed) Guidebook to Cytokines and Their receptors", Oxford Univ. Press, Oxford, pp198-201).

Each of these growth factors mediate their biological actions through binding to the corresponding receptor. The IR, IGF-1R and insulin receptorrelated receptor (IRR), for which the ligand is not known, are closely related to each other and are referred to as the insulin receptor subfamily. There is a

large body of information now available concerning the primary structure of these insulin receptor subfamily members (Ebina, Y., et al., 1985 Cell 40: 747-758; Ullrich, A., et al., 1985, Nature 313: 756-761; Ullrich, A. et al., 1986, EMBO J 5: 2503-2512; Shier, P. & Watt, V. M., 1989, J. Biol. Chem. 264: 14605-14608) and the identification of some of their functional domains (for 5 reviews see De Meyts, P. 1994, Diabetologia 37: 135-148; Lee, J. & Pilch, P. F. 1994 Amer. J. Physiol. 266: C319-C334.; Schaffer, L. 1994, Eur. J. Biochem. 221: 1127-1132). IGF-1R, IR and IRR are members of the tyrosine kinase receptor superfamily and are closely related to the epidermal growth factor receptor (EGFR) subfamily, with which they share significant sequence 10 identity in the extracellular region as well as in the cytoplasmic kinase domains (Ullrich, A. et al., 1984 Nature 309: 418-425; Ward, C. W. et al., 1995 Proteins: Structure Function & Genetics 22: 141-153). Both the insulin and EGF receptor subfamilies have a similar arrangement of two homologous domains (L1 and L2) separated by a cys-rich region of approximately 160 amino acids containing 22-24 cys residues (Bajaj, M., et al., 1987 Biochim. Biophys. Acta 916: 220-226; Ward, C. W. et al., 1995 Proteins: Structure Function & Genetics 22: 141-153). The C-terminal portion of the IGF-1R ectodomain (residues 463 to 906) is comprised of four domains: a connecting domain, two fibronectin type 3 (Fn3) repeats, and an insert domain (O'Bryan, 20 J. P., et al., 1991 Mol Cell Biol 11: 5016-5031); the C-terminal portion of the EGFR ectodomain (residues 477-621) consists solely of a second cys-rich region containing 20 cys residues (Ullrich, A. et al., 1984, Nature 309: 418-

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Little is known about the secondary, tertiary and quaternary structure of the ectodomains of these receptor subfamilies. Unlike the members of the EGFR subfamily which are transmembrane monomers which dimerise on binding ligand, the IR subfamily members are homodimers, held together by disulphide bonds. The extracellular region of the IR/IGF-1R/IRR monomers contains an  $\alpha$ -chain ( $\sim$  703 to 735 amino acid residues) and 192-196 residues of the  $\mathcal{B}$ -chain. There is a  $\sim 23$  residue transmembrane segment, followed by the cytoplasmic portion (354 to 408 amino acids) which contains the catalytic tyrosine kinase domain flanked by juxtamembrane and C-tail regulatory regions and is responsible for mediating all receptor-specific functions (White, M. F. & Kahn, C. R. 1994 J. Biol. Chem. 269: 1-4). Chemical analyses of the receptor suggest that the  $\alpha$ -chains are linked to the  $\beta$ -chains

via a single disulphide bond with the IR dimer being formed by at least two α-α disulphide linkages (Finn, F. M., et al., 1990, Proc. Natl. Acad. Sci. 87: 419-423; Chiacchia, K. B., 1991, Biochem. Biophys. Res. Commun. 176, 1178-

1182; Schaffer, L. & Ljungqvist, L., 1992, Biochem. Biophys. Res. Comm. 189: 650-653; Sparrow, L. G., et al., 1997, J. Biol. Chem. 47: 29460-29467).

Although the 3D structures of the ligands EGF, TGF-alpha (Hommel, U., et al., 1992, J. Mol. Biol. 227:271-282), insulin (Dodson, E. J., et al., 1983, Biopolymers 22:281-291), IGF-1 (Sato, A., et al., 1993, Int J Peptide Protein Res 41:433-440) and IGF-2 (Torres, A. M., et al.,1995, J. Mol. Biol. 248:385-401) are known and numerous analytical and functional studies of ligand binding to EGFR (Soler, C. & Carpenter, G., 1994 In Nicola (ed) Guidebook to Cytokines and Their receptors", Oxford Univ. Press, Oxford, pp194-197), IGF-1R and IR (see De Meyts, P., 1994 Diabetologia, 37:135-148) have been carried out, the mechanisms of ligand binding and subsequent transmembrane signalling have not been resolved.

Ligand-induced, receptor-mediated phosphorylation is the signalling mechanism by which most cytokines, polypeptide hormones and membrane-anchored ligands exert their biological effects. The primary kinase may be part of the intracellular portion of the transmembrane receptor protein as in the tyrosine kinase receptors (for review see Yarden, Y., et al., 1988, Ann. Rev. Biochem. 57:443-478) or the Ser/Thr kinase receptors (Alevizopoulos, A. & Mermod, N., 1997, BioEssays, 19:581-591) or be non-covalently associated with the cytoplasmic tail of the transmembrane protein(s) making up the receptor complex as in the case of the haemopoietic growth factor receptors (Stahl, N., et al., 1995, Science 267:1349-1353). The end result is the same, ligand binding leads to receptor dimerization or oligomerization or a conformational change in pre-existing receptor dimers or oligomers resulting in activation by transphosphorylation, of the covalently attached or non-covalently associated protein kinase domains (Hunter, T., 1995, Cell, 80:225-236).

Many oncogenes have been shown to be homologous to growth factors, growth factor receptors or molecules in the signal transduction pathways (Baserga, R.,1994 Cell, 79:927-930; Hunter, T., 1997 Cell, 88:333-346). One of the best examples is v-Erb (related to the EGFR). Since overexpression of a number of growth factor receptors results in ligand-dependent transformation an alternate strategy for oncogenes is to regulate

the expression of growth factor receptors or their ligands or to directly bind to the receptors to stimulate the same effect (Baserga, R., 1994 Cell, 79:927-930). Examples are v-Src, which activates IGF-1 R intracellularly; c-Myb, which transforms cells by enhancing the expression of IGF1R and SV40 T antigen which interacts with the IGF-1R and enhances the secretion of IGF-1 (see Baserga, R.,1994 Cell, 79:927-930 for review). Cells in which the IGF-1 receptor has been knocked out cannot be transformed by SV40 T antigen. If oncogenes activate growth factors and their receptors then tumour suppressor genes should have the opposite effect. One good example of this is WT1, the Wilm's tumour suppressor gene which suppresses the expression of IGF-1R (Drummond, J. A., et al., 1992, Science, 257:275-277). Cells that are driven to proliferate by oncogenes undergo massive apotosis when growth factor receptors are ablated since unlike normal cells, they appear unable to withdraw from the cell-cycle and enter into the G0 phase (Baserga, R.,1994 Cell, 79:927-930).

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The insulin-like growth factor-1 receptor (IGF-1R) is one of several growth-factor receptors that regulate the proliferation of mammalian cells. However, its ubiquitousness and certain unique aspects of its function make IGF-1R an ideal target for therapeutic interventions against abnormal growth, with very little effect on normal cells (see Baserga, R., 1996 TIBTECH, 14:150-152). The receptor is activated by IGF1, IGF2 and insulin and plays a major role in cellular proliferation in at least three ways: it is essential for optimal growth of cells in vitro and in vivo; several cell types require IGF-1R to maintain the transformed state and activated IGF-1R has a protective effect against apoptotic cell death (Baserga, R., 1996 TIBTECH, 14:150-152). These properties alone make it an ideal target for therapeutic interventions. Transgenic experiments have shown that IGF-1R is not an absolute requirement for cell growth but is essential for the establishment of the transformed state (Baserga, R., 1994 Cell, 79: 927-930). In several cases (human glioblastoma, human melanoma; human breast carcinoma; human lung carcinoma; human ovaraian carcinoma; human rhabdomyosarcoma; mouse melanoma, mouse leukaemia; rat glioblastoma; rat rhabdomyosarcoma; hamster mesothelioma) the transformed phenotype can be reversed by decreasing the expression of IGF-1R using antisense to IGF-1R (Baserga, R., 1996 TIBTECH 14:150-152); or interfering with its function by antibodies to IGF-1R (human breast carcinoma; human rhabdomyosarcoma)

or by dominant negatives of IGF-1R (rat glioblastoma; Baserga, R.,1996 TIBTECH 14:150-152).

Three effects are observed when the function of IGF-1R is impaired:

tumour cells undergo massive apoptosis which results in inhibition of tumourogenesis; surviving tumour cells are eliminated by a specific immune 5 response; and such a host response can cause a regression of an established wild-type tumour (Resnicoff, M., et al., 1995, Cancer Res. 54:2218-2222). These effects, plus the fact that interference of IGF-1R function has a limited effect on normal cells (partial inhibition of growth without apoptosis) makes IGF-1R a unique target for therapeutic interventions (Baserga, R., 1996 10 TIBTECH 14:150-152). In addition IGF-1R is downstream of many other growth factor receptors, which makes it an even more generalised target. The implication of these findings is that if you can decrease the number of IGF-1 receptors on cells or antagonise their function then tumours cease to grow and can be removed immunologically. These studies establish that IGF-1R 15 antagonists will be extremely important therapeutically.

Many cancer cells have constitutively active EGFR (Sandgreen, E. P., et al., 1990, Cell, 61:1121-135; Karnes, W. E. J., et al., 1992, Gastroenterology, 102:474-485) or other EGFR family members (Hines, N. E.,1993, Semin. Cancer Biol. 4:19-26). Elevated levels of activated EGFR occur in bladder, breast, lung and brain tumours (Harris, A. L., et al., 1989, In Furth & Greaves (eds) The Molecular Diagnostics of human cancer. Cold Spring Harbor Lab. Press, CSH, NY, pp353-357). Antibodies to EGFR can inhibit ligand activation of EGFR (Sato, J. D., et al., 1983 Mol. Biol. Med. 1:511-529) and the growth of many epithelial cell lines (Aboud-Pirak E., et al., 1988, J. Natl Cancer Inst. 85:1327-1331). Patients receiving repeated doses of a humanised chimeric anti-EGFR antibody showed signs of disease stabilization. The large doses required and the cost of production of humanised Mab is likely to limit the application of this type of therapy. These findings indicate that the development of EGF antagonists will be attractive anticancer agents.

## Summary of the Invention

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The present inventors have now obtained 3D structural information concerning the insulin-like growth factor receptor (IGF-1R) and the insulin receptor (IR) which provides a rational basis for the development of antagonists and agonists of the polypeptide ligands for specific therapeutic applications. This information can be used to predict the structure of related

members of the insulin receptor family and epidermal growth factor family and to develop agonists and antagonists of their respective polypeptide ligands.

Accordingly, in a first apsect the present invention provides a method of screening for, or designing, an agonist of a ligand of an insulin receptor family member or EGF receptor family member which method includes

- (i) selecting or designing a substance which possesses stereochemical complementarity to a receptor site, wherein the receptor site is characterised by
- (a) amino acids 1-462 of IGF-1R positioned at atomic coordinates substantially as shown in Figure 1 or a subset thereof; or
- (b) amino acids derived from an insulin receptor family member or EGF receptor family member which form an equivalent structure to the amino acids defined in paragraph (a); and
- (ii) testing the substance for the ability to act as an agonist of the ligand of an insulin receptor family member or EGF receptor family member.

In a second apsect the present invention provides a method of screening for, or designing, an antagonist of a ligand of an insulin receptor family member or EGF receptor family member which method includes

- (i) selecting or designing a substance which possesses stereochemical complementarity to a receptor site, wherein the receptor site is characterised by
- (a) amino acids 1-462 of IGF-1R positioned at atomic coordinates substantially as shown in Figure 1 or a subset thereof; or
- (b) amino acids derived from an insulin receptor family member or an EGF receptor family member which form an equivalent structure to the amino acids defined in paragraph (a); and
- (ii) testing the substance for the ability to act as an antagonist of the ligand of an insulin receptor family member or EGF receptor family member.

The phrase "insulin receptor family" encompasses, for example, IGF-1R, IR and IRR. The phrase "EGF receptor family" encompasses for example, EGFR, ErbB2, ErbB3 and ErbB4. In general, insulin receptor family members and EGF receptor family members show similar domain arrangements and share significant sequence identity (preferably at least 20% identity between the families and at least 40% identity within each family).

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The receptor site defined in the first and second aspects of the present invention comprises the L1-cysteine rich-L2 region (residues 1-462) of the ectodomain of IGF-1R. At the centre of this structure is a groove, bounded by all three domains, of sufficient size to accommodate a ligand molecule. By "stereochemical complementarity" we mean that the biologically active substance or a portion thereof correlates, in the manner of the classic "lock-and-key" visualisation of ligand-receptor interaction, with the groove in the receptor site. Preferably, the stereochemical complementarity is such that the compound has a K<sub>I</sub> for the receptor site of less than 10<sup>-6</sup>M. More preferably, the K<sub>I</sub> value is less than 10<sup>-8</sup>M and more preferably less than 10<sup>-9</sup>M.

In preferred embodiments of the first and second aspects of the present invention, the method further involves selecting or designing a substance which has portions that match residues positioned on the surface of the receptor site which faces the groove. By "match" we mean that the identified portions interact with the surface residues, for example, via hydrogen bonding or by enthalpy-reducing Van der Waals interactions which promote desolvation of the biologically active substance within the site, in such a way that retention of the biologically active substance within the groove is favoured energetically.

In a preferred embodiment of the first aspect of the present invention, the method includes screening for, or designing, a substance which possesses a stereochemistry and/or geometry which allows it to interact with both the L1 and L2 domains of the receptor site. As described above, the insulin receptor exists as homodimers held together by disulphide bonds. Electron miscroscopy studies described herein indicate that the insulin receptor monomers dimerise in nature in such a manner that the grooves of each monomer may face each other. Accordingly, the method of the first aspect of the present invention may involve screening for, or designing, a biologically active substance which interacts with the L1 domain of one monomer and the L2 domain of the other monomer.

In a third aspect the present invention provides a method of selecting or designing an agonist of a ligand of an insulin receptor family member or EGF receptor family member which method includes

(i) selecting or designing a substance which interacts with

(a) a fragment of IGF-1R characterised by amino acids 1-462 positioned at atomic coordinates substantially as shown in Figure 1 or a subset thereof; or

(b) a fragment derived from an insulin family receptor member or EGF receptor family member which is equivalent to the fragment defined in paragraph (a);

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wherein the interaction of the substance with the fragment alters the position of at least one of the L1, L2 or cys-rich domains of the fragment relative to the position of at least one of the other domains; and

(ii) testing the substance for the ability to act as an agonist of the ligand of an insulin receptor family member or EGF receptor family member.

In a preferred embodiment of the third aspect of the present invention the substance interacts with the fragment in the region of the L1 domain-cys rich domain interface, causing the L1 and cys-rich domains to move away from each other. In a further preferred embodiment the substance interacts with the hinge region between the L2 domain and the cys-rich domain causing an alteration in the positions of the domains relative to each other. In a further preferred embodiment the substance interacts with the beta sheet of the L1 domain causing an alteration in the position of the L1 domain relative to the position of the cys-rich domain or L2 domain.

In a fourth aspect the present invention provides an agonist of a ligand of an insulin receptor family member or EGF receptor family member obtained by a method according to the first or third aspects of the present invention.

In a fifth aspect the present invention provides an antagonist of ligand of an insulin receptor family member or EGF receptor family member obtained by a method according to the second aspect of the present invention.

The agonists or antagonists of the fourth and fifth aspects of the present invention may be mutant insulin family member or EGF family member ligands where at least one mutation occurs in the region of the ligand which interacts with residues on the surface of the receptor site facing toward the groove. For example, the IGF-1 ligand has a predominance of basic residues in the C region which may interact with the acidic patch of the cys-rich region near L1. An acidic patch on the other side of the ligand may interact with the patch of basic residues (residues 307-310) on the N-terminal

end of L2. Accordingly, mutants of IGF-1 which exhibit altered activity may be generated by introducing modifications in the C region of IGF-1 or residues in the acidic patch on the other side of the hormone.

In a sixth aspect the present invention provides a substance which possesses stereochemical complementarity to a receptor site, wherein the receptor site is characterised by

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- (a) amino acids 1-462 of IGF-1R positioned at atomic coordinates substantially as shown in Figure 1 or a subset thereof; or
- (b) amino acids derived from an insulin receptor family member or an EGF receptor family member which form an equivalent structure to the amino acids defined in paragraph (a);

with the proviso that the substance is not a naturally occurring ligand of an insulin receptor family member or EGF receptor family member or a mutant thereof.

By "mutant" we mean a ligand which has been modified by one or more point mutations, insertions of amino acids or deletions of amino acids.

In a preferred embodiment of the sixth aspect of the present invention, the stereochemical complementarity is such that the compound has a  $K_I$  for the receptor site of less than  $10^{-6}M$ . More preferably, the  $K_I$  value is less than  $10^{-8}M$  and more preferably less than  $10^{-9}M$ .

In a seventh aspect the present invention provides a pharmaceutical composition for treatment of a disease associated with reduced activity of a ligand of an insulin receptor family member or EGF receptor family member which includes an agonist obtained by a method according to the first or third aspects of the present invention and a pharmaceutically acceptable carrier or diluent.

In an eighth aspect the present invention provides a pharmaceutical composition for treatment of a disease associated with activity of a ligand of an insulin receptor family member or EGF receptor family member which includes an antagonist obtained by a method according to the second aspect of the present invention and a pharmaceutically acceptable carrier or diluent.

In a ninth aspect the present invention provides a method of preventing or treating a disease associated with reduced activity of a ligand of an insulin receptor family member or EGF receptor family member which method includes administering to a subject in need thereof an agonist obtained by a method according to the first or third aspects of the present invention.

Diseases associated with reduced activity of a ligand of an insulin receptor family member or EGF receptor family member include diabetes, osteoporosis, nerve degeneration and a range of catabolic states.

In a tenth aspect the present invention provides a method of preventing or treating a disease associated with activity of a ligand of an insulin receptor family member or EGF receptor family member which method includes administering to a subject in need thereof an antagonist obtained by a method according to the second aspect of the present invention.

Diseases associated with activity of a ligand of an insulin receptor family member or EGF receptor family member include cancer, leukaemia and many types of tumour states including but not restricted to breast cancer, brain tumours, ovarian cancer, pancreatic tumours, lung cancer, melanoma, rhabdomyosarcoma, mesothelioma and glioblastoma.

## **Brief Description of the Drawings**

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Figure 1. IGF-1R residues 1-462, in terms of atomic coordinates refined to a resolution of 2.6 Å (average accuracy  $\approx$  0.3Å). The coordinates are in relation to a Cartesian system of orthogonal axes.

Figure 2. Depiction of the residues lining the groove of the IGF-1R receptor fragment 1-462.

Figure 3. Gel filtration chromatography of affinity-purified IGF-1R/462 protein. The protein was purified on a Superdex S200 column (Pharmacia) fitted to a BioLogic L.C. system (Biorad), equilibrated and eluted at 0.8 ml/min with 40 mM Tris/150 mM NaCl/0.02% NaN3 adjusted to pH 8.0. (a) Protein eluting in peak 1 contained aggregated IGF-1R/462 protein, peak 2 contained monomeric protein and peak 3 contained the c-myc undecapeptide used for elution from the Mab 9E10 immunoaffinity column. (b) Non-reduced SDS-PAGE of fraction 2 from IGF-1R/462 obtained following Superdex S200 (Fig.1a). Standard proteins are indicated.

Figure 4. Ion exchange chromatography of affinity-purified, truncated IGF-1R ectodomain. A mixture of gradient and isocratic elution chromatography was performed on a Resource Q column (Pharmacia) fitted to a BioLogic System (Biorad), using 20 mM Tris/pH 8.0 as buffer A and the same buffer containing 1M NaCl as buffer B. Protein solution in TBSA was diluted at least 5 1:2 with water and loaded onto the column at 2 ml/min. Elution was monitored by absorbance (280 nm) and conductivity (mS/cm). Target protein (peak 2) eluted isocratically with 20 mM Tris/0.14 M NaCl pH 8.0. Inset: Isoelectric focusing gel (pH 3 - 7; Novex Australia Pty Ltd)of fraction 2. The pI was estimated at 5.1 from standard proteins (not shown). 10

Figure 5. Gel filtration chromatography of affinity purified IR/485 protein. Affinity-purified material at 1 mg/ml produced a dominant peak at apparent mass ~ 140 kDa (interpreted as a dimer) (a); whereas affinity-purified material at 0.02 mg/ml produced a dominant peak at apparent mass ~ 85kDa (interpreted as a monomer) (b).

Figure 6. (a) SDS-PAGE of IR/485 following gel filtration chromatography. The protein migrated as a single broad band of apparent molecular mass ~ 78 kDa (reduced - lane A) or ~ 68kDa (non-reduced - lane B). (b) Isoelectric 20 focussing of the IR/485 protein. The IR/485 fragment reacted positively in an ELISA with Mab 83-7, gave a single sequence corresponding to the Nterminal 10 residues of IR, showing several isoforms on isoelectric focussing from pI6.0-6.8. The fragment was further purified by ion-exchange chromatography on Uno Q (BioRad, USA), using stepwise isocratic elution with incremental changes in salt concentrations (see Figure 7). Fractions A and D were each enriched in a component isoform from the ladder of isoforms present in the unfractionated mixture. Both these fractions produced crystals, whereas no crystals were obtained from fractions B and C.

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Figure 7. Purification of the IR/485 protein by ion-exchange chromatography on Uno Q (BioRad, USA), using stepwise isocratic elution with incremental changes in salt concentrations.

35 Figure 8. Polypeptide fold for residues 1-462 of IGF-1R. The L1 domain is at the top, viewed from the N-terminal end and L2 is at the bottom. The space

at the centre is of sufficient size to accommodate IGF-1. Helices are indicated by curled ribbon and b-strands by arrows. Cysteine side chains are drawn as ball-and-stick with lines showing disulfide bonds. The arrow points in the direction of view for Figure 9.

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Figure 9. Amino acid sequences of IGF-1R and related proteins. a, L1 and L2 domains of human IGF-1R and IR are shown based on a sequence alignment for the two proteins and a structural alignment for the L1 and L2domains. Positions showing conservation physico-chemical properties of amino acids are boxed, residues used in the structural alignment are shaded yellow and residues which form the Trp 176 pocket are in red. Secondary structure elements for L1 (above the sequences) and L2 (below) are indicated as cylinders for helices and arrows for b-strands. Strands are colour coded according to the b-sheet to which they belong. Disulfide bonds are also indicated. b, Cys-rich domains of human IGF-1R, IR and EGFR (domains 2 and 4) are aligned based on sequence and structural considerations. Secondary structural elements and disulfide bonds are indicated above the sequences. The dashed bond is only present in IR. Different types of disulfide bonded modules are labelled below the sequences as open, filled or broken lines. Boxed residues show conservation of physico-chemical properties and structurally conserved residues for modules 4-7 are shaded yellow. Residues from EGFR which do not conform to the pattern are shaded grey and the conserved Trp 176 and the semi-conserved Gln 182 are shaded red. This figure was prepared using ALSCRIPT (Barton, G. J., 1993, Prot. Engineering, 6:37-40).

Figure 10. Stereo view of a superposition of the L1 (white) and L2 (black) domains. Residues numbers above are for L1 and below for L2. The side chain of Trp 176 which protrudes into the core of L1 is drawn as ball-and-stick.

- Figure 11. Schematic diagram showing the association of three  $\beta$ -finger motifs.  $\beta$ -strands are drawn as arrows and disulfide bonds as zigzags.
- Figure 12. GRASP [Nicolls, A. et al., 1993, Biophys. J. 64, 166-170] surface diagram of the L1 domain of IGF-1R shown in a similar view to Figure 8. The

N-terminal β-strand is at the top. The mutation L87A [ Nakae, J. et al., 1995, J. Biol. Chem. 270, 22017-22022] and four regions (residues 12-15, 34-44, 64-67 and 89-91 of IR) shown to be important in insulin binding to IR [Williams, P. F. et al., 1995, J. Biol. Chem. 270, 3012-3016] correspond to a patch of residues on the large β-sheet. Residues numbers for IR/IGF-1R are given and residues are coloured according to the magnitude of Kd(mutant)/Kd(wild type), red, > 40; orange, 10-40; yellow, 2.5-10; green, < 2.5; non-secreting, white; untested, blue. All mutants on the opposite face of the domain do not affect insulin affinity.

Figure 13: Sequence Alignment of hIGF-1R, hIR and hIRR Ectodomains. Derived by use of the PileUp program in the software package of the Genetics Computer Group, 575 Science Drive, Madison, Wisconsin, USA. For assignment of homologous 3D structures see Figure 9.

Figure 14: Sequence Alignment of EGFR, ErbB2, ErbB3 and ErbB4 Ectodomains. Derived by use of the PileUp program in the software package of the Genetics Computer Group, 575 Science Drive, Madison, Wisconsin, USA. For alignment on the IGF-1R fragment and assignment of homologous 3D structures, see Figure 9.

Figure 15 Sequence Alignment and Classification of the Disulphide-bonded Modules in the Cys-rich domains of IGF-1R, IR, IRR, EGFR, ErbB2, ErbB3 and ErbB4.

Figure 16. Gel filtration chromatography of insulin receptor ectodomain and MFab complexes. hIR -11 ectodomain dimer (5 - 20 mg) was complexed with MFab derivatives (15-25 mg each) of the anti-hIR antibodies 18-44, 83-7 and 83-14 (Soos et al., 1986). Elution profiles were generated from samples loaded onto a Superdex S200 column (Pharmacia), connected to a BioLogic chromatography system (Biorad) and monitored at 280 nm. The column was eluted at 0.8 ml/min with 40 mM Tris/150 mM sodium chloride/0.02% sodium azide buffer adjusted to pH 8.0: Profile 0, hIR -11ectodomain, Profile 1, ectodomain mixed with MFab 18-44; Profile 2, ectodomain mixed with MFab 18-44 and MFab 83-14; Profile 3, ectodomain mixed with MFab 18-44, MFab 83-14 and MFab 83-7. The apparent mass of each complex was

determined from a plot of the following standard proteins: thyroglobulin (660 kDa), ferritin (440 kDa), bovine gammaglobulin (158 kDa), bovine serum albumin (67 kDa), chicken ovalbumin (44 kDa) and equine myoglobin (17 kDa).

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- Figure 17. Micrographs of hIR and hIGF-1R ectodomains.(a) Undecorated hIR ectodomain dimer stained with methylamine tungstate showing parallel bars. (b) Undecorated hIR ectodomain dimer stained with uranyl formate, showing well-spaced parallel bars corresponding to the cartoon below.

  (c) Undecorated hIGF-1R ectodomain dimer stained with uranyl formate.

  Magnification bars for (a), (b) and (c) 50nm.
- Figure 18. Micrographs of hIR and hIGF-1R ectodomains. (a) Thinly stained region of undecorated hIR ectodomain dimers in uranyl formate, showing U-shaped particles (circled) as well as parallel bars as in the cartoon below. (b) Undecorated hIGF-1R ectodomain dimer under similar staining conditions. Magnification bars 50 nm.
- Figure 19. hIR ectodomain dimer complexed with MFab 83-7 and stained with KPT. Three projections can be recognised: circled particles have the Fab arms displaced either clockwise as in the cartoon below left,or anticlockwise as in the cartoon below middle; arrowed particles have the Fab arms in a central position, cartoon below right. Magnification bar 50 nm.
- Figure 20. hIR ectodomain dimer complexed with MFab 83-7 and stained with uranyl formate showing the parallel bar structure in particles having the Fab arms displaced (circled). Magnification bar 50 nm.
- Figure 21. (a) hIR ectodomain dimer complexed with MFab 83-14 stained with potassium phosphotungstate, showing Fab arms attached near the bottom of U-shaped particles (circled). The corresponding cartoon is shown below left. (b) hIR ectodomain dimer complexed with MFab 83-14 stained with uranyl acetate, showing both the view described above (circled) and the parallel-bar view with diagonally projecting Fab arms (arrowed), as in the cartoon below right. Magnification bars 50 nm.

Figure 22. Double complex of hIR ectodomain dimer with MFabs 83-7 and 18-44 showing particles of complex shape (circled) with four Fab arms attached, consistent with the cartoon below. Magnification bar 50 nm.

- Figure 23. Images of hIR ectodomain dimer co-complexed with MFabs 83-7, 83-14 and 18-44 showing examples of complex particles (circled) where it is possible to identify that there are more than four MFabs bound to the dimeric central region. Magnification bar 50 nm.
- Figure 24. Schematic illustrating the proposed model of the hIR ectodomain dimer. The dimensions of the molecular envelope are as shown in the diagram, as is the position of the two-fold axis.

## **Detailed Description of the Invention**

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We describe herein the expression, purification, and crystallization of a recombinant IGF-1R fragment (residues 1-462) containing the L1-cysteinerich-L2 region of the ectodomain. The selected truncation position is just downstream of the exon 6/exon 7 junction (Abbott, A. M., et al., 1992. J Biol Chem., 267:10759-10763) and occurs at a position where the sequences of the IR and EGFR families diverge markedly (Ward, C. W., et al., 1995, Proteins: Struct., Funct., Genet. 22:141-153; Lax, I., et al., 1988, Molec. Cellul. Biol. 8:1970-1978) suggesting it represents a domain boundary. To limit the effects of glycosylation, the IGF-1R fragment was expressed in Lec8 cells, a glycosylation mutant of Chinese hamster ovary (CHO) cells, whose defined glycosylation defect produces N-linked oligosaccharides truncated at N-acetyl glucosamine residues distal to mannose residues (Stanley, P. 1989, Molec. Cellul. Biol. 9:377-383). Such an approach has facilitated glycoprotein crystallization (Davis, S. J., et al., 1993, Protein Eng. 6:229-232; Liu, J., et al., 1996, J. Biol. Chem. 271:33639-33646).

The IGF-1R construct described herein included a c-myc peptide tag (Hoogenboom, H. R., et al.,1991, Nucleic Acids Res. 19:4133-4137) that is recognised by the Mab 9E10 (Evan, G. I., et al., 1985, Mol. Cell. Biol. 5:3610-3616) enabling the expressed product to be purified by peptide elution from an antibody affinity column followed by gel filtration over Superdex S200.

The purified proteins crystallized under a sparse matrix server (Israerile I. 8)

The purified proteins crystallized under a sparse matrix screen (Jancarik, J. & Kim, S.-H., 1991, J. Appl. Cryst. 24:409-411) but the crystals were of variable

quality, with the best diffracting to 3.0-3.5Å. Isocratic gradient elution by anion-exchange chromatography yielded protein that was less heterogenous and gave crystals of sufficient quality to determine the structure of the first three domains of the human IGF-1R.

The IGF-1R fragment consisted of residues 1-462 of IGF-1R linked via an enterokinase-cleavable pentapeptide sequence to an eleven residue c-myc peptide tag at the C-terminal end. The fragment was expressed in Lec8 cells by continuous media perfusion in a bioreactor using porous carrier disks. It was secreted into the culture medium and purified by peptide elution from an anti-c-myc antibody column followed by Superdex S200 gel filtration. The receptor fragment bound two anti-IGF-1R monoclonal antibodies, 24-31 and 24-60, which recognize conformational epitopes, but could not be shown to bind IGF-1 or IGF-2. Crystals of variable quality were grown as rhombic prisms in 1.7 M ammonium sulfate at pH 7.5 with the best diffracting to 3.0-3.5 Å. Further purification by isocratic elution on an anion-exchange column gave protein which produced better quality crystals, diffracting to 2.6 Å, that were suitable for X-ray structure determination.

The structure of this fragment (IGF-1R residues 1-462; L1-cys rich-L2domains) has been determined to 2.6 Å resolution by X-ray diffraction. The L domains each adopt a compact shape consisting of a single stranded right-handed β-helix. The cys-rich region is composed of eight disulphide-bonded modules, seven of which form a rod-shaped domain with modules associated in a novel manner. At the centre of this reasonably extended structure is a space, bounded by all three domains, and of sufficient size to accommodate a ligand molecule. Functional studies on IGF-1R and other members of the insulin receptor family show that the regions primarily responsible for hormone-binding map to this central site. Thus this structure gives a first view of how members of the insulin receptor family might interact with their ligands.

Another group has reported the crystallization of a related receptor, the EGFR in a complex with its ligand EGF (Weber, W., et al., 1994, J Chromat. 679:181-189). However difficulties were encountered with these crystals which diffracted to only 6 Å, insufficient for the determination of an atomic resolution structure of this complex (Weber, W., et al., 1994, J Chromat 679:181-189) or the generation of accurate models of structurally related receptor domains such as IGF-1R and IR by homology modelling.

The present inventors have applied the same process to the IR and generated a fragment (residues 1-485) that covers the first three domains of the IR. This fragment has been expressed in transformed Lec8 cells, purified, and crystallized by similar methodologies to yield crystals suitable for X-ray diffraction.

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The present inventors have therefore developed 3D structural information about cytokine receptors to enable a more accurate understanding of how the binding of ligand leads to signal transduction. Such information provides a rational basis for the development of antagonists or agonists for specific therapeutic applications, something that heretofore could not have been predicted *de novo* from available sequence data.

The precise mechanisms underlying the binding of agonists and antagonists to the IGF-1 receptor site are not fully clarified. However, the binding of the agonists or antagonists to the receptor site, preferably with an affinity in the order of 10<sup>-8</sup>M or higher, is understood to arise from enhanced stereochemical complementarity, relative to naturally occurring IGF-1 ligands.

Such stereochemical complementarity, pursuant to the present invention, is characteristic of a molecule that matches intra-site surface residues lining the groove of the receptor site as eneumerated by the coordinates set out in Figure 1. The residues lining the groove are depicted in Figure 2. Substances which are complemetary to the shape of the receptor site characterised by amino acids positioned at atomic coordinates set out in Figure 1 may be able to bind to the receptor site and, when the binding is sufficiently strong, substantially prohibit binding of the naturally occurring ligands to the site.

It will be appreciated that it is not necessary that the complementarity between agonists or antagonists and the receptor site extend over all residues lining the groove in order to inhibit binding of the natural ligand. Accordingly, agonists or antagonists which bind to a portion of the residues lining the groove are encompassed by the present invention.

In general, the design of a molecule possessing stereochemical complementarity can be accomplished by means of techniques that optimize, either chemically or geometrically, the "fit" between a molecule and a target receptor. Known techniques of this sort are reviewed by Sheridan and Venkataraghavan, Acc. Chem Res. 1987 20 322; Goodford, J. Med. Chem.

1984 <u>27</u> 557; Beddell, Chem. Soc. Reviews 1985, 279; Hol, Angew. Chem. 1986 <u>25</u> 767 and Verlinde C.L.M.J & Hol, W.G.J. Structure 1994, <u>2</u>, 577, the respective contents of which are hereby incorporated by reference. See also Blundell et al., Nature 1987 <u>326</u> 347 (drug development based on information regarding receptor structure).

Thus, there are two preferred approaches to designing a molecule, according to the present invention, that complements the shape of IGF-1R or a related receptor molecule. By the geometric approach, the number of internal degrees of freedom (and the corresponding local minima in the molecular conformation space) is reduced by considering only the geometric (hard-sphere) interactions of two rigid bodies, where one body (the active site) contains "pockets" or "grooves" that form binding sites for the second body (the complementing molecule, as ligand). The second preferred approach entails an assessment of the interaction of respective chemical groups ("probes") with the active site at sample positions within and around the site, resulting in an array of energy values from which three-dimensional contour surfaces at selected energy levels can be generated.

The geometric approach is illustrated by Kuntz et al., J. Mol. Biol. 1982 161 269, the contents of which are hereby incorporated by reference, whose algorithm for ligand design is implemented in a commercial software package distributed by the Regents of the University of California and further described in a document, provided by the distributor, which is entitled "Overview of the DOCK Package, Version 1.0,", the contents of which are hereby incorporated by reference. Pursuant to the Kuntz algorithm, the shape of the cavity represented by the IGF-R1 site is defined as a series of overlapping spheres of different radii. One or more extant data bases of crystallographic data, such as the Cambridge Structural Database System maintained by Cambridge University (University Chemical Laboratory, Lensfield Road, Cambridge CB2 1EW, U.K.) and the Protein Data Bank maintained by Brookhaven National Laboratory (Chemistry Dept. Upton, NY 11973, U.S.A.), is then searched for molecules which approximate the shape thus defined.

Molecules identified in this way, on the basis of geometric parameters, can then be modified to satisfy criteria associated with chemical complementarity, such as hydrogen bonding, ionic interactions and Van der Waals interactions.

The chemical-probe approach to ligand design is described, for example, by Goodford, J. Med. Chem. 1985 <u>28</u> 849, the contents of which are hereby incorporated by reference, and is implemented in several commercial software packages, such as GRID (product of Molecular Discovery Ltd., West

- Way House, Elms Parade, Oxford OX2 9LL, U.K.). pursuant to this approach, the chemical prerequisites for a site-complementing molecule are identified at the outset, by probing the active site (as represented via the atomic coordinates shown in Fig. 1) with different chemical probes, e.g., water, a methyl group, an amine nitrogen, a carboxyl oxygen, and a hydroxyl.
- Favored sites for interaction between the active site and each probe are thus determined, and from the resulting three-dimensional pattern of such sites a putative complementary molecule can be generated.

The chemical-probe approach is especially useful in defining variants of a molecule known to bind the target receptor. Accordingly, crystallographic analysis of IGF-1 bound to the receptor site may provide useful information regarding the interaction between the archetype ligand and the active site of interest.

In summary, the general principles of receptor-based drug design can be applied by persons skilled in the art, using the crystallographic results presented above, to produce agonists or antagonists of IGF-1R having sufficient stereochemical complementarity to exhibit high affinity binding to the receptor site.

The present invention is further described below with reference to the following, non-limiting examples.

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## **EXAMPLE 1**

## Expression, Purification and Crystalization of the IGF-1R Fragment.

Several factors hamper macromolecular crystallization including sample selection, purity, stability, solubility (McPherson, A., et al., 1995, Structure 3:759-768); Gilliland, G. L., & Ladner, J. E., 1996, Curr. Opin. Struct. Biol. 6:595-603), and the nature and extent of glycosylation (Davis, S. J., et al., 1993, Protein Eng. 6:229-232). Initial attempts to obtain structural data from soluble IGF-1R ectodomain (residues 1-906) protein, expressed in Lec8 cells (Stanley, P. 1989, Molec. Cellul. Biol. 9:377-383) and purified by affinity chromatography, produced large, well-formed crystals (1.0 mm x 0.2 mm) which gave no discernable X-ray diffraction pattern

(unpublished data). Similar difficulties have been encountered with crystals of the structurally related epidermal growth factor receptor (EGFR) ectodomain which diffracted to only 6 Å, insufficient for the determination of an atomic resolution structure (Weber, W. et al., 1994, J Chromat 679:181-189). This prompted us to search for a fragment of IGF-1R that was more amenable to X-ray crystallographic studies.

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The fragment expressed (residues 1-462) comprises the L1-cysteinerich-L2 region of the ectodomain. The selected truncation position at Val462 is four residues downstream of the exon 6/exon 7 junction (Abbott, A. M., et al., 1992, J Biol Chem. 267:10759-10763) and occurs at a position where the sequences of the IR and the structurally related EGFR families diverge markedly (Lax, I., et al., 1988, Molec Cell Biol. 8:1970-1978; Ward, C. W., et al., 1995, Proteins: Struct., Funct., Genet. 22:141-153), suggesting it represents a domain boundary. The expression strategy included use of the pEE14 vector (Bebbington, C. R. & Hentschel, C. C. G., 1987, In: Glover, D. M., ed. DNA Cloning. Academic Press, San Diego. Vol 3, p163) in glycosidase-defective Lec8 cells (Stanley, P., 1989, Molec. Cellul. Biol. 9:377-383), which produce N-linked oligosaccharides lacking the terminal galactose and N-acetylneuraminic acid residues (Davis, S. J., et al., 1993, Protein Eng. 6:229-232; Liu, T., et al., 1996, J Biol Chem 271:33639-33646.). The construct contained a C-terminal c-myc affinity tag (Hoogenboom, H. R., et al., 1991, Nucl Acids Res. 19:4133-4137), which facilitated immunoaffinity purification by specific peptide elution and avoided aggressive purification conditions. These procedures yielded protein which readily crystallized after a gel filtration polish. This provided a general protocol to enhance crystallisation prospects for labile, multidomain glycoproteins.

The structure of this fragment is of considerable interest since it contains the major determinants governing insulin and IGF-1 binding specificity (Gustafson, T. A. & Rutter, W. J., 1990, J. Biol. Chem. 265:18663-18667; Andersen, A. S., et al., 1990, Biochemistry, 29:7363-7366; Schumacher, R., et al., 1991, J. Biol. Chem. 266:19288-19295; Schumacher, R., et al., 1993, J. Biol. Chem. 268:1087-1094; Schäffer, L., et al., 1993, J. Biol. Chem. 268:3044-3047; Williams, P. F., et al., 1995, , J. Biol. Chem. 270:3012-3016) and is very similar to an IGF-1R fragment (residues 1-486) reported to act as a strong dominant negative for several growth functions and which

induces apoptosis of tumour cells in vivo (D'Ambrosio, C., et al., 1996, Cancer Res. 56:4013-4020).

The expression plasmid pEE14/IGF-1R/462 was constructed by inserting the oligonucleotide cassette:

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**AatII** 

5' GACGTC GACGATGACGATAAG GAACAAAAACTCATC

D V D D D D K E Q K L I (EK cleavage) (c-myc tail)

10 S E E D L N (Stop)

TCAGAAGAGGATCTGAAT TAGAATTC GACGTC 3'

EcoRI AatII

encoding an enterokinase cleavage site, c-myc epitope tag (Hoogenboom, H. R., et al., 1991, Nucleic acids Res. 19:4133-4137) and stop codon into the AatII site (within codon 462) of IGF-1 receptor cDNA in the mammalian expression vector pECE (Ebina, Y., et al., 1985, Cell, 40:747-758; kindly supplied by W. J. Rutter, UCSF, USA), and introducing the DNA comprising the 5' 1521 bp of the cDNA (Ullrich, A., et al., 1986, EMBO J. 5:2503-2512) ligated to the oligonucleotide cassette into the EcoRI site of the mammalian plasmid expression vector pEE14 (Bebbington, C. R. & Hentschel, C. C. G., 1987, In: Glover, D. M., ed. DNA Cloning. Academic Press, San Diego. Vol 3, p163; Celltech Ltd., UK). Plasmid pEE14/IGF-1R/462 was transfected into Lec8 mutant CHO cells (Stanley, P. 1989, Molec. Cellul. Biol. 9:377-383) obtained from the American Tissue Culture Collection (CRL:1737) using Lipofectin (Gibco-BRL). Cell lines were maintained after transfection in glutamine-free medium (Glascow modification of Eagle's medium (GMEM; ICN Biomedicals, Australia) and 10% dialysed FCS (Sigma, Australia) containing 25  $\mu$ M methionine sulphoximine (MSX; Sigma, Australia) as described (Bebbington, C. R. & Hentschel, C. C. G., 1987, In: Glover, D. M., ed. DNA Cloning. Academic Press, San Diego. Vol 3, p163). Transfectants were screened for protein expression by Western blotting and sandwich enzyme-linked immunosorbant assay (ELISA) (Cosgrove, L., et al., 1995, ) using monoclonal antibody (Mab) 9E10 (Evan et al., 1985) as the capture antibody and either biotinylated anti-IGF-1R Mab 24-60 or 24-31 for detection(Soos et al., 1992; gifts from Ken Siddle, University of Cambridge,

UK). Large-scale cultivation of selected clones expressing IGF-1R/462 was carried out in a Celligen Plus bioreactor (New Brunswick Scientific, USA) containing 70 g Fibra-Cel Disks (Sterilin, UK) as carriers in a 1.25 L working volume. Continuous perfusion culture using GMEM medium supplemented with non-essential amino acids, nucleosides, 25  $\mu M$  MSX and 10% FCS was maintained for 1 to 2 weeks followed by the more enriched DMEM/F12 without glutamine, with the same supplemention for the next 4-5 weeks. The fermentation production run was carried out three times under similar conditions and resulted in an estimated overall yield of 50 mg of receptor protein from 430 L of harvested medium. Cell growth was poor during the initial stages of the fermentation when GMEM medium was employed, but improved dramatically following the switch to the more enriched medium. Target protein productivity was essentially constant during the period from ~100 to 700 h of the 760 h fermentation, as measured by ELISA using Mab 9E10 as the capture antibody and biotinylated Mab 24-31 as the developing antibody.

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Soluble IGF-1R/462 protein was recovered from harvested fermentation medium by affinity chromatography on columns prepared by coupling Mab 9E10 to divinyl sulphone-activated agarose beads (Mini Leak; Kem En Tec, Denmark) as recommended by the manufacturer. Mini-Leak Low and Medium affinity columns with antibody loadings of 1.5-4.5 mg/ml of hydrated matrix were obtained, with the loading range of 2.5-3 mg/ml giving optimal performance (data not shown). Mab 9E10 was produced by growing hybridoma cells (American Tissue Culture Collection) in serum-free medium in the Celligen Plus bioreactor and recovering the secreted antibody (4 g) using protein A glass beads (Prosep-A, Bioprocessing Limited, USA). Harvested culture medium containing IGF-1R/462 protein was adjusted to pH 8.0 with Tris-HCl (Sigma), made 0.02% (w/v) in sodium azide and passed at 3-5 ml/min over 50 ml Mab 9E10 antibody columns at 4° C. Bound protein was recovered by recycling a solution of 2-10 mg of the undecamer c-myc peptide EQKLISEEDLN (Hoogenboom et al., 1991) in 20 ml of Tris-buffered saline containing 0.02% sodium azide (TBSA). Between 65% and 75% of the product was recovered from the medium as estimated by ELISA, with a further 15-25% being recovered by a second pass over the columns. Peptide recirculation (~10 times) through the column eluted bound protein more efficiently than a single, slower elution. Residual bound protein was eluted

with sodium citrate buffer at pH 3.0 into 1 M Tris HCl pH 8.0 to neutralize the eluant, and columns were re-equilibrated with TBSA.

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Gel filtration over Superdex S200 (Pharmacia, Sweden), of affinitypurified material showed a dominant protein peak at ~63 kDa, together with a smaller quantity of aggregated protein (Figure 3a). The peak protein migrated primarily as two closely spaced bands on reduced, sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE; Figure 3b), reacted positively in the ELISA with both Mab 24-60 and Mab 24-31, and gave a single sequence corresponding to the N-terminal 14 residues of IGF-1R. No binding of IGF-1 or IGF-2 could be detected in the solid plate binding assay (Cosgrove et al., 1995, Protein Express Purif. 6:789-798). The IGF-1R/462 fragment was further purified by ion-exchange chromatography on Resource Q (Pharmacia, Sweden). Using shallow salt gradients, protein enriched in the slowest migrating SDS-PAGE band was obtained (data not shown), which formed relatively large, well-formed crystals (see below). Isoelectric focusing showed the presence of one major and two minor isoforms. Protein purified on Resource Q with an isocratic elution step of 0.14 M NaCl in 20 mM TrisCl at pH 8.0 (fraction 2, Figure 4) showed less heterogeneity on isoelectric focusing (Figure 4 inset) and SDS-PAGE (data not shown) and produced crystals of sufficient quality for structure determination (see below).

Crystals were grown by the hanging drop vapour diffusion method using purified protein concentrated in Centricon 10 concentrators (Amicon Inc, USA) to 5-10 mg/ml in 10-20 mM Tris-HCl pH 8.0 and 0.02% (w/v) azide, or 100 mM ammonium sulfate and 0.02% (w/v) azide. A search for crystallization conditions was performed initially using the factorial screen (Jancarik, J. & Kim, S.-H.,1991, J Appl Cryst 24:409-411) and subsequently optimised. Crystals were examined on an M18XHF rotating anode generator (Siemens, Germany) equipped with Franks mirrors (MSC, USA) and RAXIS IIC and IV image plate detectors (Rigaku, Japan).

From the initial crystallization screen of this protein, crystals of about 0.1 mm in size grew in one week. Upon refining conditions, crystals of up to 0.6 x 0.4 x 0.4 mm could be grown from a solution of 1.7-2.0 M ammonium sulfate, 0.1 M HEPES pH 7.5. The crystals varied considerably in shape and diffraction quality, growing predominantly as rhombic prisms with a length to width ratio of up to 5:1, but sometimes as rhombic bipyramids, the latter form being favoured when using material which had been eluted

from the Mab 9E10 column at pH 3.0. Each crystal showed a minor imperfection in the form of very faint lines from the centre to the vertices. Protein from dissolved crystals did not appear to be different from the protein stock solution when run on an isoelectric focusing gel. Upon X-ray examination, the crystals diffracted to 3.0-4.0 Å and were found to belong to the space group  $P2_12_12_1$  with a = 76.8 Å, b = 99.0 Å, c = 119.6 Å. In the diffraction pattern, the crystal variability noted above was manifest as a large (1-2°) and anisotropic mosaic spread, with concomitant variation in resolution. To improve the quality of the crystals, they were grown in the presence of various additives or were recrystallized. These methods failed to substantially improve the crystal quality although bigger crystals were obtained by recrystallization. The variability in crystal quality appeared to be due to protein heterogeneity, as demonstrated by the observation that more highly purified protein, eluted isocratically from the Resource Q column and showing one major band on isoelectric focusing (Figure 4 inset), produced crystals of sufficient quality for structure determination. These crystals diffracted to 2.6 Å resolution with cell dimensions, a = 77.0 Å, b = 99.5 Å, c = 120.1 Å and mosaic spread of 0.5°. Heavy metal derivatives of the IGF-1R/462 crystals have been obtained and are leading to the determination of an atomic resolution structure of this fragment, which contains the L1, cysteine-rich and L2 domains of human IGF-1R.

## **EXAMPLE 2**

## **Expression, Purification and Crystalization of the IR Fragment**

A similar strategy was adopted for the human insulin receptor. The fragment expressed (residues 1-485) comprises the L1-cysteine-rich-L2 region of the IR ectodomain but extends 13 residues further before the attachment of the 17 residue EK cleavage site linker and c-myc tail. The selected truncation position corresponds to a unique and convenient Bgl II restriction site. The expression strategy was also based on the pEE14 expression vector in glycosidase-defective Lec8 cells and use of a C-terminal c-myc affinity tag for immunoaffinity purification by specific peptide elution. These procedures yielded IR protein which readily crystallized after a gel filtration polish.

The expression plasmid pHIR485 was constructed by ligating the double-stranded oligonucleotide cassette:

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Bgl II Xba I

#### 5' AGATC TCCGACGATGACGATAAG GAACAAAAACTCATCTCAGAAGAGGGATCTGAAT TAG TCTAGA 3'

#### KISDDDDKEQKLISEEDLN

EK cleavage c-myc tail Stop

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encoding an enterokinase cleavage site, c-myc epitope tag (Hoogenboom, H. R., et al., 1991, Nucleic acids Res. 19:4133-4137) and stop codon, to the larger 11.1 kilobasepair Bgl II / Xba I fragment isolated from digestion of the mammalian expression plasmid pEH3 (a derivative of the mammalian plasmid expression vector pEE14 [Bebbington, C. R. & Hentschel, C. C. G., 1987, In: Glover, D. M., ed. DNA Cloning. Academic Press, San Diego. Vol 3, p163; Celltech Ltd., UK] which holds the entire coding sequence of human insulin receptor within a Hind III /Xba I fragment). Lec8 mutant CHO cells (Stanley, P. 1989, Molec. Cellul. Biol. 9:377-383) obtained from the American Tissue Culture Collection (CRL:1737) were transfected with pHIR485 using Lipofectamine (Gibco-BRL). Cell lines were maintained after transfection in glutamine-free medium (Glascow modification of Eagle's medium - GMEM; ICN Biomedicals, Australia) and 10% dialysed FCS (Sigma, Australia) containing 25 µM methionine sulphoximine (MSX; Sigma, Australia) as described (Bebbington, C. R. & Hentschel, C. C. G., 1987, In: Glover, D. M., ed. DNA Cloning. Academic Press, San Diego. Vol 3, p163). Transfectants were screened for protein expression by Western blotting and sandwich enzyme-linked immunosorbant assay (ELISA) (Cosgrove, L., et al., 1995, ) using anti-hIR (Mab) 83.7 as the primary antibody and biotinylated monoclonal antibody (Mab) 9E10 (Evan et al., 1985) for detection (Soos et al., 1986; gifts from Ken Siddle, University of Cambridge, UK). Large-scale cultivation of selected clones expressing IR/485 was carried out in a Celligen Plus bioreactor (New Brunswick Scientific, USA) containing 70 g Fibra-Cel Disks (Sterilin, UK) as carriers in a 1.25 L working volume. Continuous perfusion culture was carried out using DMEM/F12 without glutamine medium (ICN), supplemented with non-essential amino acids, nucleosides, 25 µM MSX and 5 - 10% FCS and resulted in an estimated overall yield of 115 mg of receptor protein from 165 L of harvested medium. Target protein productivity was essentially constant during the fermentation, as measured by ELISA.

Soluble IR/485 protein was recovered from harvested fermentation medium by affinity chromatography on columns of Mab 9E10 essentially as described in Example 1. Between 92 -98% of the product was recovered from the medium by this affinity-chromatography step, as estimated by ELISA.

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Gel filtration over Superdex 200 (Pharmacia, Sweden), of the affinitypurified material at 1mg/ml produced a dominant protein peak at apparent mass ~140 kDa (Figure 5a - interpreted as dimer), whereas a peak at apparent mass ~85 kDa was obtained (Figure 5b - interpreted as monomer) at 0.02 mg/ml. The protein migrated as a single broad band of apparent molecular mass ~78 kDa (reduced- lane A) or ~68 kDa (non-reduced - lane B) on sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE; Figure 6a) The IR/485 fragment reacted positively in the ELISA with Mab 83-7, gave a single sequence corresponding to the N-terminal 10 residues of IR, showing several isoforms on isoelectric focussing from pI 6.0 - 6.8 (Figure 6b). Crystallisation screening trials of the fragment produced crystals too small for X-ray diffraction studies. The fragment was further purified by ionexchange chromatography on Uno Q (BioRad, USA), using stepwise isocratic elution with incremental changes in salt concentrations (Figure 7). Fractions A and D were each enriched in a component isoform from the ladder of isoforms present in the unfractionated mixture (Figure 6b). Both these fractions produced crystals, whereas no crystals were obtained from fractions B and C.

Crystals were grown by the hanging drop vapour diffusion method using purified protein concentrated in Centricon 10 concentrators (Amicon Inc, USA) to 5-10 mg/ml in 10mM Tris-HCl pH 8.0 and 0.02% (w/v) azide. A search for crystallization conditions was performed initially using the factorial screen (Jancarik, J. & Kim, S.-H.,1991, J Appl Cryst 24:409-411) and subsequently optimised. Crystals were examined on an M18XHF rotating anode generator (Siemens, Germany) equipped with Franks mirrors (MSC, USA) and an RAXIS IIC image plate detector (Rigaku, Japan).

From the initial crystallization screen of this protein fraction D fine needles grew in about one week. In further experiments, crystals of up to  $0.04 \times 0.04 \times 0.2$  mm could be grown from a solution of 1.9-2.0 M ammonium sulfate, 2% PEG 400, 0.1 M HEPES pH 7.5. Upon X-ray examination, the crystals diffracted to 4 Å and were found to belong to the space group  $P2_12_12_1$  with a = 103.2 Å, b = 130.0 Å, c = 161.6 Å. Despite their small size these

crystals diffracted sufficiently well to allow collection of a low resolution data set. Further purification of the protein and refinement of crystallisation conditions should yield larger crystals, providing data to determine the structure of this fragment at medium resolution or better.

## 5 EXAMPLE 3

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## Structure of the IGF-1R/1-462

Crystals were cryo-cooled to-170°C in a mother liquor containing 20% glycerol, 2.2 M ammonium sulfate and 100 mM Tris at pH 8.0. Native and derivative diffraction data were recorded on Rigaku RAXIS IIc or IV area detectors using copper Kα radiation from a Siemens rotating anode generator with Yale/MSC mirroroptics. The space group was P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub> with a = 77.39 Å, b = 99.72 Å, and c = 120.29 Å. Data were reduced using DENZO and SCALEPACK (Otwinowski, Z. & Minor, W., 1996, Mode.Meth. Enzym. 276:307-326). Diffraction was notably anisotropic for all crystals examined.

Phasing by multiple isomorphous replacement(MIR) was performed with PROTEIN (Steigeman, W. Dissertation (Technical Univ. Munich, 1974) using anomalous scattering for both UO2 and PIP derivatives. Statistics for data collection and phasing are given in Table 1. In the initial MIR map regions of protein and solvent could clearly be seen but the path of the polypeptide was by no means obvious. That map was subject to solvent flattening and histogram matching in DM (Cowtan, K., 1994, Joint CCP4 and ESF-EACBM newslett. Protein Crystallogr. 31:34-38). The structure was traced and rebuilt using O (Jones, T. A., et al., 1991, Acta Crystallogr. A47:110-119) and refined with X-PLOR 3.851 (Brunger, A. T., 1996, X-PLOR ReferenceManual 3.851, Yale Univ., New Haven, CT). After 5 rounds of rebuilding and energy minimisation the R-factor dropped to 0.279 and Rfree = 0.359 for data 7-2.6 Å resolution. The current model contains 458 amino acids and 3 N-linked carbohydrates but no solvent molecules. For residues with B(Ca) > 70 Å2atomic positions are less reliable (37-42, 155-159, 305, 336-341, 404-406,453-458). There is weak electron density for residues 459-461 but the c-myc tail appears completely disordered.

The 1-462 fragment consists of the N-terminal three domains of IGF-1R (L1, cys-rich, L2) and contains regions of the molecule which dictate ligand specificity (17-23). The molecule adopts a reasonably extended structure (approximately  $40 \times 48 \times 105 \text{ Å}$ ) with domain 2 (cys-rich region) making contact along the length of domain 1 (L1) but very little contact with

the third domain (L2) (see Figure 8). This leaves a space at the centre of the molecule of approximately  $24 \text{ Å} \times 24 \text{ Å} \times 24 \text{ Å}$  which is bounded on three sides by the three domains of the molecule. The space is of sufficient size to accommodate the ligand; IGF-1.

### The L domains

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Each of the L domains (residues 1-150 and 300-460) adopt a compact shape (24 x 32 x 37 Å) consisting of a single-stranded right handed  $\beta$ -helix and capped on the ends by short a-helices and disulfide bonds. The body of the domain looks like a loaf of bread with the base formed from a flat sixstranded  $\beta$ -sheet, 5 residues long and the sides being  $\beta$ -sheets three residues long (Figures 8 & 9). The top is irregular but in places is similar for the two domains. The two domains are superposable with an rms deviation in Ca positions of 1.6 Å for 109 atoms (Figure 10). Although this fold is reminiscent of other  $\beta$ -helix proteins it is much simpler and smaller with very few elaborations and thus it represents a new superfamily of domains. One notable difference between the two domains is that the indole ring of Trp 176 from the cys-rich region (Figure 9b) is inserted into the hydrophobic core of L1 and the C-terminal helix is only vestigial (Figure 8). For the insulin receptor family the sequence motif of residues which form the Trp pocket in L1 does not occur in L2 (Figure 9a). However in the EGF receptor, which has an additional cys-rich region after the L2 domain (14, 15), the pocket motif can be found in both L domains and the Trp is conserved in both cys-rich regions (Figure 9b).

The repetitive nature of the β-helix is reflected in the sequence and the first five turns were correctly identified by Bajaj, M., et al. (1987, Biochim.Biophys. Acta 916:220-226), the conserved Gly residues being found in turns making one bottom edge of the domain. However, their conclusions about the fold were incorrect. The"helix-like" repeat is actually a pair of bends at the top edge of the domain. In their Motif V, the Gly is not in a bend but is followed by the insertion of a conserved loop of 7-8 residues (see Figure 9a). Glycine is structurally important in the Gly bends as mutation of these residues compromises folding of the receptor [van der Vorm, E.R., et al., 1992, J. Biol. Chem. 267, 66-71; Wertheimer, E. et al., 1994, J. Biol. Chem. 269, 7587-7592].

Upon comparing the L domains with other right-handed β-helix structures such as pectate lyase (Yoder, M. D., et al., 1993,.Structure, 1:241-

251-1507) and the p22 tailspike protein (Steinbacher, S., et al., 1997, J.Mol. Biol. 267:865-880) there are some striking similarities as well as differences. In all cases the ends of the domain are capped by α-helices but the L domains also have a disulphide bond at each end to hold the termini. The other β-helix domains are considerably longer and have significant twist to their sheets while the L domains have flat sheets. Although the sizes of the helix repeats are similar (here 24-25 residues vs 22-23 for pectate lyase) the cross-sections are quite different. The L domains have a rectangular cross-section while pectate lyase and p22 tailspike protein are V-shaped and have many, and sometimes quite large, insertions (Yoder, M. D., et al., 1993, Structure, 1:241-251-1507; Steinbacher, S., et al., 1997, J.Mol. Biol. 267:865-880). In the hydrophobic core a common feature is the stacking of aliphatic residues from successive turns of the β-helix and near the C-terminus of each L domain there is also a short Asn ladder, reminiscent of the long Asn ladder observed in pectate lyase (Yoder, M. D., et al., 1993, Structure 1:241-251-

1507). On the opposite side of the L domains the Gly bend as well as the two

domains. Thus although the L domains are built on similar principles to the

bends and sheet preceding it have no counterpart in the other  $\beta$ -helix

other  $\beta$ -helix domains they constitute a separate superfamily.

## 20 The cys-rich domain

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The cys-rich domain is composed of eight disulfide-bonded modules (Figure 9b), the first of which sits at the end of L1 while the remainder make a curved rod running diagonally across L1 and reaching to L2 (Figure 8). The strands in modules 2-7 run roughly perpendicular to the axis of the rod in a manner more akin to laminin (Stetefeld, J., et al., 1996, J.Mol. Biol. 257:644-657) than to TNF receptor (Banner, D. W., et al., 1993, Cell, 73:431-445) but the modular arrangement of the cys-rich domain is different to other cys-rich proteins for which structures are known. The first 3 modules of IGF-1R have a common core, containing a pair of disulfide bonds, but show considerable variation in the loops (Figure 9b). The connectivity of these modules is the same as the first half of EGF (Cys 1-3and 2-4) but their structures do not appear to be closely related to any member of the EGF family. Modules 4 to 7 have a different motif,  $\beta$ -finger, and best match residues 2152-2168 of fibrillin (Dowling, A. K., et al., 1996, Cell, 85:597-605). Each is composed of three polypeptide strands, the first and third being disulfide bonded and the latter two forming a  $\beta$ -ribbon. The  $\beta$ -ribbon of each  $\beta$ - finger module lines up antiparallel to form a tightly twisted 8-stranded  $\beta$ -sheet (Figures 8 and 11). Module 6 deviates from the common pattern with the first segment being replaced by an  $\alpha$ -helix followed by a large loop that is likely to have a role in ligand binding (see below). As module 5 is most similar to module 7 it is possible that the four modules arose from serial gene duplications. The final module is a disulfide linked bend of five residues.

The fact that the two major types of cys-rich modules occur separately implies that these are the minimal building blocks of cys-rich domains found in many proteins. Although it can be as short as 16 residues, the motif of modules 4-7 is clearly distinct and capable of forming a regular extended structure. Thus cys-rich domains such as these can be considered as made of repeat units each composed of a small number of modules.

## Hormone binding

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Attempts have been made to locate the IGF-1 (and insulin) binding site by examining natural (Taylor, S. I., 1992, Diabetes, 41:1473-1490) and **15** site-directed mutants (Williams, P. F., et al., 1995, J. Biol. Chem. 270:3012-3016; Mynarcik, D. C et al., 1996, J. Biol. Chem. 271:2439-2442; Mynarcik, D. C., et al., 1997, J. Biol. Chem. 272:2077-2081), chimeric receptors (Andersen, A. S., et al., 1990, Biochemistry 29:7363-7366; Gustafson, T. A., & Rutter, W. J., 1990, J. Biol. Chem. 265:18663-18667; Schäffer, L., et al., 1993, J. Biol. 20 Chem. 268:3044-3047; Schumacher, R., 1993, J. Biol. Chem. 268:1087-1094; Kjeldsen, T., et al., 1991, Proc. Natl Acad. Sci. USA, 88:4404-4408) and by crosslinking studies (Wedekind, F., et al., 1989, Biol. Chem Hoppe-Seyler, 370:251-258; Fabry, M., 1992, J. Biol. Chem. 267:8950-8956; Waugh, S. M., et al., 1989, Biochemistry, 28:3448-3458; Kurose, T., et al., 1994),.J. Biol. 25 Chem.269:29190-29197-34). IGF-1R/IR chimeras not only show which regions of the receptors account for ligand specificity but also provide an efficient means of identifying some parts of the hormone binding site. Paradoxically regions controlling specificity are not the same for insulin and IGF-1. Replacing the first 68 residues of IGF-1R with those of IR confers 30 insulin binding ability on the chimeric IGF-1R (Kjeldsen, T., et al., 1991, Proc. Natl Acad. Sci. USA, 88:4404-4408) and replacing residues 198-300 in the cys-rich region of IR with the corresponding residues 191-290 of IGF-1R allows the chimeric receptor to bind IGF-1 (Schäffer, L., et al.,1993, J. Biol. Chem. 268:3044-3047). Thus a receptor can be constructed which binds both 35

IGF-1 and insulin with near native affinity. From the structure it is clear that if the hormone bound in the central space it could contact both these regions.

From analysis a series of chimeras examined by Gustafson, T. A., & Rutter, W. J. (J. Biol. Chem. 265:18663-18667, 1990) the specificity determinant in the cys-rich region can be limited further to residues 223-274. 5 This region corresponds to modules 4-6 and includes a large and somewhat mobile loop (residues 255-263, mean B[Ca atoms] = 57 Å2) which extends into the central space (see Figure 8). In IR this loop is four residues bigger and is stabilised by an additional disulfide bond (Schäffer, L. & Hansen, P.H.,1996, Exp. Clin. Endocrinol. Diabetes, 104: Suppl. 2, 89). The larger 10 loop of IR may serve to exclude IGF-1 from the hormone binding site but allow the smaller insulin molecule to bind. It is interesting to note that mosquito IR homologue, which has a loop two residues larger than the mammalian IRs, also appears to bind insulin but not IGF-1 (Graf, R., et al., 1997, Insect Molec.Biol. 6:151-163). Analysis of the structure indicates that 15 the insulin/IGF-1 specificity is controlled by residues in this loop (amino

acids 253-272 in IGF-1R; amino acids 260-283 in IR)

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As chimeras only address residues which differ between the two receptors a more precise analysis of the site can be obtained from single site mutants. In particular, from an alanine-replacement study, four regions of L1 important for insulin binding were identified (Williams, P. F., et al., 1995, J. Biol. Chem. 270:3012-3016). The first three are at similar positions on successive turns of the b-helix and the fourth lies on the conserved bulge on the large b-sheet (Figure 12). Thus there is a footprint for insulin binding to the L1 domain which lies on the first half of large b-sheet facing into the central space. Residues further along the sheet which are conserved in IGF-1R and could also be important. The conservative substitution of leucine for methionine at residue 119 of IR (113 of IGF-1R) causes a mild form of leprechaunism [Hone, J. et al., 1994, J. Med. Genet. 31, 715-716]. This residue is buried and the mutation could perturb neighbouring residues to affect insulin binding.

The axis of the L2 domain is perpendicular to that of the L1 domain and N-terminal end of its  $\beta$ -helix is presented to the hormone-binding site. On this face of the L2 domain the only mutation studied so far is the naturally occurring IR mutant, S323L, which gives rise to Rabson-Mendehall syndrome and severe insulin resistance (Roach, P.,1994, Diabetes 43:1096-

1102). As this mutant only affects insulin binding and not cell-surface expression, residue 323 of IR (residue 313 of IGF-1R) is probably at or near the binding site. Structurally this residue lies in the middle of a region (residues 309-318 of IGF-1R) which is conserved in both IR and IGF-1R and the surrounding region, 332-345 (of IGF-1R), is also quite well conserved in the these receptors (Figure 9a). Therefore this region is quite likely to form part of the hormone-binding site but would not have been detected by chimeras. It is interesting to note that in this region IRR is not as well conserved as the other two receptors (Shier, P. & Watt, V.M., 1989, J.Biol.Chem. 264:4605-14608).

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The distance from this putative hormone-binding region on L2 to that found on L1 is about 30 Å (Figure 8). Thus L1 and L2 appear too far apart to bind IGF-1 or insulin. However, in the crystal structure there is a deep cleft between part of the cys-rich domain (residue 262)and L2 (residue 305) and this cleft is occupied by a loop from a neighbouring molecule. Thus it seems probable that the position of the L2 domain in the receptor structure or the hormone-receptor complex adopts a different position with respect to the cys-rich domain than that found in the crystal. The movement required to bring L2 sufficiently close to L1 is small, namely a rotation of approximately 25° about residue 298.

A number of IR mutants have been identified which constitutively activate the receptor and the majority of these are found in the  $\alpha$  chain. Curiously all  $\alpha$  chain mutants involve changes to or from proline or the deletion of an amino acid, implying that they cause local structural rearrangements. The mutation R86N is similar to wild type but R86P reduces cell-surface expression and insulin binding while constitutively activating autophosphorylation [Grønskov, K. et al., 1993, Biochem. Biophys. Res. Commun. 192, 905-911]. The proline mutation probably disturbs residues preceding 87 which lie in the interface between the L1 and cys-rich domains but it could also affect insulin binding. In the cys-rich domain residues 233, 281, 244 and 247 of IR are not conserved in IGF-1R (Figure 9b) yet L233P [Klinkhamer, M.P. et al., 1989, EMBO J. 8, 2503-2507], deletion of N281 [Debois-Mouthon, C. et al., 1996, J. Clin. Endochronol. Metab. 81, 719-727] or the triple mutant P243R, P244R and H247D [Rafaeloff, R. et al., 1989, J. Biol. Chem. 264, 15900-15904] cause constitutive kinase activation. Due to their locations each of these three mutants appears likely to compromise the

folding of a β-finger domain and, in turn, the structural integrity of the rodlike cys-rich domain. The structural ramifications of these mutations could be significant for the whole receptor ectodomain as disturbing the L1/cys-rich interface or distorting the rod-like domain could affect the relative position of L1 and the cys-rich domain in this context.

L1 has been further implicated as deletion of K121 on the opposite side of L1 from the cys-rich domain was also found to cause autophosphorylation [Jospe, N. et al., 1994, J. Clin. Endochronol. Metab. 79, 1294-1302]. By contrast this mutation does not affect insulin binding. Thus a possible mechanism emerges for insulin binding and signal transduction. When insulin binds between L1 and L2 it modifies the relative position of L1 and the cys-rich domain in the receptor, perhaps by hinge motion between L2 and the cys-rich domain like that suggested above, and the structural rearrangement is transmitted across the plasma membrane. In the absence of insulin the same signal can be initiated by mutations in the cys-rich region or at the L1/cys-rich interface but at the expense on insulin binding. The signal can also be initiated more directly by mutations on the opposite side of L1 which affect the interaction of L1 with other parts of the ectodomain, possibly the other half of the receptor dimer.

## Ligand Studies

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Although there is no structural information about an IGF-1/IGF-1R complex a number of studies have probed the nature of this interaction. Results from cross-linking experiments with IGF-1 and insulin and their cognate receptors are consistent with the hormone binding site proposed above. For example B29 of insulin can be cross-linked to the cys-rich region (residues 205-316( (Yip, C. C., et al., 1988, Biochim. Biophys. Res. Commun. 157:321-329) or the L1 domain (Wedekind, F., et al., 1989, Biol. Chem Hoppe-Seyler, 370:251-258). However these two regions are reasonably well separated and those studies may indicate that B29 is mobile. Other studies unfortunately do not map the site any more precisely.

Analogues and site-directed mutants of IGF-1 and -2 have been more fruitful. Relative to insulin IGF-1 and -2 contain two extra regions, the C region between B and A and a D peptide at the C-terminus. For IGF-1 replacement of the C region by a four Gly linker reduced affinity for IGF-1R by a factor of 40 but increased affinity for IR 5-fold (Bayne, M.L.,et al., 1988, J. Biol.Chem. 264:11004-11008). Changes in affinity are consistent with the

deletion in IGF-1 complementing differences in the cys-rich regions of IGF-1R and IR noted above. Mutation of residues either side of the C region (residue 24 for IGF-1 [Cascieri, M.A., et al., 1988, Biochemistry 27:3229-323], residues 27,43 for IGF-2, [Sakano, K., et al., 1991, J. Biol. Chem.

266:20626-20635]) also have deleterious effects on the affinity of the hormone forIGF-1R as has truncation of the nearby D peptide in IGF-2 (Roth, B.V., et al., 1991, Biochem. Biophys. Res. Commun. 181:907-914). Insulin has been extensively mutated. Binding studies [summarised in Kristensen, C. et al., 1997, J. Biol. Chem. 272, 12978-12983] indicate that insulin may bind its receptor via a hydrophobic patch (residues A2, A3, A19, B8, B11, B12, B15 and possibly B23 & B24). However this patch is normally buried and requires the removal of the B chain's C-terminus from the observed position. Assuming IGF-1, -2 and insulin bind their receptors in the same orientation, these data suggest an approximate orientation for the hormone when bound to the receptor.

One notable feature of IGF-1 and -2 is the large number of charged residues and their uneven distribution over the surface. Basic residues are predominantly found in the C region and, in solution, this region is not well ordered in either IGF-1 or -2 (Sato, A., et al., 1993, Int J Peptide Protein Res. 41:433-440; Torres, A. M., et al., 1995, J. Mol. Biol. 248:385-401). In contrast the binding site of the receptor has a sizable patch of acidic residues in the corner where the cys-rich domain departs from L1. Other acidic residues which are specific to this receptor are found along the inside face of the cys-rich domain and the loop (residues 255-263) extending from module 6. Thus it is possible that electrostatics play an important part in IGF-1 binding with the C region binding to the acidic patch of the cys-rich region near L1 and the acidic patch on the other side of the hormone directed towards a small patch of basic residues (residues 307-310) on the N-terminal end of L2.

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Although the structure of this fragment gives significant information about the nature of the hormone binding site, residues outside this region have also been shown to affect binding of ligand. A number of studies have implicated residues 704-715 of IR (Mynarcik, D. C et al., 1996, J. Biol. Chem. 271, 2439-2442; Kurose, T., et al., 1994, J. Biol. Chem.269:29190-29197). These residues could contact insulin on one of the sides left open in the current structure. Using insulin labelled at the B1 residue, Fabry, M., et al., (1992, J. Biol. Chem. 267:8950-8956) cross linked insulin to the fragment

390-488, part of which is not near the site as described. The explanation for this could be either 488 reaches back to the hormone binding site, or this region could contact another hormone bound to the other half of the receptor.

Further structural information is needed to establish how these other regions contact the hormone and to elucidate how binding of the hormone is communicated to the kinase inside the cell.

The structure of the L1-cys-rich-L2 domains of IGF-1R presented here represents the first structural information for the extracellular portion of a member of the insulin receptor family. The L domains display a novel fold which is common to the EGF receptor family and the modular architecture of the cys-rich domain implies that smaller building blocks should be used to describe the composition of cysteine-rich domains. This fragment contains the major specificity determinants of receptors of this class for their ligands. It has an elongated structure with a space in the middle which could accommodate the ligand. The three sides of this site correspond to regions which have been implicated in hormone binding. Although other sites are present in the receptor ectodomain which interact with the ligand this structure gives us an initial view of how the insulin, IGF-1 and -2 might interact with their cell surface receptors to control their metabolic and mitogenic effects

Such information will provide valuable insight into the structure of the corresponding domains of the IR and insulin receptor-related receptor as well as members of the related EGFR family (Bajaj, M., et al., 1987, Biochim Biophys Acta 916:220-226; Ward, C. W. et al., 1995, Proteins: Struct Funct Genet 22:141-153).

#### **EXAMPLE 4**

### <u>Prediction of 3D Structure of the Corresponding Domains of IRR and IR</u> <u>Based on Structure of IGF-1R Frgament.</u>

The sequence identities between the different members of the insulin receptor family are sufficient to allow accurate sequence alignments to facilitate 3D structure predictions by homology modelling. The alignments of the ectodomains of human IGF-1R, IR, and IRR are shown in Figure 13.

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#### **EXAMPLE 5**

# Prediction of 3D Structure of EGFR and its Family Members ERB2, ERB3 and ERB4.

The sequence identities between the different members of the EGFR receptor family and the insulin receptor family are sufficient to allow accurate sequence alignments to facilitate 3D structure predictions by homology modelling. The alignments of the ectodomains of human EGFR, ERB2, ERB3 and ERB4 are shown in Figure 14. The ectodomains of the EGFR family members are composed of four domains: L1 domain, cys-rich domain, L2 domain and a second cys-rich domain all of which can be modelled from the structure of the IGF-1R fragment residues 1-462.

The sequence alignment analysis and characterization of the repeat modules in the cys-rich region of IGF-1R and the homologous regions of the IR, IRR and the first and second cys-rich regions of EGFR, ErbB2, ErbB3 and ErbB4 are shown in Figure 15. A representative of each subtype of cys repeat is found in the IGF-1R fragment 1-462 and is used to model each of these modules in the other receptors. Note the nature and order of modules in the second cys-rich repeat of the EGFR family is different to that seen in the first cys-rich region.

#### **EXAMPLE 6**

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# Single-Molecule Imaging of Human Insulin Receptor Ectodomain and its Fab Complexes

### Cloning and expression of hIR -11 ectodomain protein

A full length clone of the human IR exon -11 form (hIR -11) was prepared by exchanging an Aat II fragment, nucleotides 1195 to 2987, of the 25 exon +11 clone (plasmid pET; Ellis et al., 1986; gift from Dr W. J. Rutter, UCSF) of hIR (Ebina et al., 1985, Cell 40, 747-758) with the equivalent Aat II fragment from a plasmid (pHIR/P12-1, ATCC 57493) encoding part of the extracellular domain and the entire cytoplasmic domain of hIR -11 (Ullrich et al., 1985, Nature 313, 756-761). The ectodomain fragment of hIR -11 30 (2901 bp, coding for the 27 residue signal sequence and residues His1-Asn914) was produced by SalI and SspI digestion and inserted into the mammalian expression vector pEE6.HCMV-GS (Celltech Limited, Slough, Berkshire, UK) into which a stop codon linker had been inserted, as described previously (Cosgrove et al., 1995, Protein Expression and 35 Purification 6, 789-798) for the hIR exon +11 ectodomain.

The resulting recombinant plasmid pHIR II (2 µg) was transfected into glycosylation deficient Chinese hamster ovary (Lec 8) cells (Stanley, 1989, Molec. Cellul. Biol. 9, 377-383) with Lipofectin (Gibco-BRL). After

transfection, the cells were maintained in glutamine-free medium GMEM (ICN Biomedicals, Australia) as described previously (Bebbington & Hentschel, 1987, In DNA Cloning (Glover, D., ectodomain.), Vol III, Academic Press, san Diego; Cosgrove et al., 1995, Protein Expression and Purification 6, 789-798). Expressing cell lines were selected for growth in GMEM with 25 µM methionine sulphoximine (MSX, Sigma). Transfectants were screened for protein expression using sandwich ELISA with anti-IR monoclonal antibodies 83-7 and 83-14. Metabolic labelling of cells, immunoprecipitations, insulin binding assays and Scatchard analyses were performed as described previously for the exon +11 form of hIR ectodomain (Cosgrove et al., 1995, , Protein Expression and Purification 6, 789-798).

#### hIR -11 ectodomain production and purification

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The selected clone (inoculum of 1.28 x 108 cells) was grown in a spinner flask packed with 10 g of Fibra-cel disc carriers (Sterilin, U.K.) in 500 ml of GMEM medium containing 10% fetal calf serum (FCS) and 25  $\mu$ M MSX. Selection pressure was maintained for the duration of the culture.

Ectodomain was recovered from harvested media by affinity chromatography on immobilized insulin and further purified by gel filtration chromatography on Superdex S200 (Pharmacia; 1 x 40 cm) in Tris-buffered saline containing 0.02% sodium azide (TBSA) as described previously (Cosgrove et al., 1995, *Protein Expression and Purification* 6, 789-798). Solutions of purified hIR -11 ectodomain were stored at 4° C prior to use.

### Production of Fab fragments and their complexes with ectodomain

Purification of Mabs 83-7, 83-14 and 18-44 from ascites fluid by affinity chromatography using Protein A-Sepharose, and the production of Fabs, were based on the methodologies described in Coligan et al.,1993, Current Protocols in Immunology, Vol 1, pp 2.7.1-2.8.9, Greene Publishing Associates & Wiley - Interscience, John Wiley and Sons. Fab was produced from monoclonal antibody by mercuripapain digestion for 1-4 h, followed by gel filtration on Superdex S200. Products were monitored by reducing and non-reducing SDS-PAGE. For 83-7 Mab, an IgG Type 1 monoclonal antibody, the bivalent (Fab)2' isolated by this method was reduced to monovalent Fab 83-7 by mild reduction with mM L-cysteine.HCl in 100 mM Tris pH 8.0

(Coligan et al., 1993, Current Protocols in Immunology, Vol 1, pp 2.7.1-2.8.9, Greene Publishing Associates & Wiley - Interscience, John Wiley and Sons).

Complexes of Fab with hIR -11 ectodomain were produced by mixing

~ 2.5 to 3.5 molar excess of Fab with hIR -11 ectodomain at ambient temperature in TBSA at pH 8.0. After 1-3 h, the complex was separated from unbound Fab by gel filtration over a Superdex S200 column in the same buffer.

#### **Electron microscopy**

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Uncomplexed hIR -11 ectodomain and the Fab complexes described above were diluted in phosphate-buffered saline (PBS) to concentrations of the order of 0.01-0.03 mg/ml. Prior to dilution, 10% glutaraldehyde (Fluka) was added to the PBS to achieve a final concentration of 1% glutaraldehyde. Droplets of ~ 3ml of this solution were applied to thin carbon film on 700mesh gold grids after glow-discharging in nitrogen for 30 s. After 1 min. the excess protein solution was drawn off and followed by application and withdrawal of 4-5 droplets of negative stain [2% uranyl acetate (Agar), 2% uranyl formate (K and K), 2% potassium phosphotungstate (Probing and Structure) adjusted to pH 6.0 with KOH, or 2% methylamine tungstate (Agar) adjusted to pH 6.8 with NH4OH]. In the case of both uranyl acetate and uranyl formate staining, an intermediate wash with 2 or 3 droplets of PBS was included prior to application of the stain. The grids were air-dried and then examined at 60kV accelerating voltage in a JEOL 100B transmission electron microscope at a magnification of 100,000x. It was found that there was a typical thickness of negative stain in which Fabs were most easily seen, hence areas for photography had to be chosen from particular zones of the grid. Electron micrographs were recorded on Kodak SO-163 film and developed in undiluted Kodak D19 developer. The electron-optical magnification was calibrated under identical imaging conditions by recording single-molecule images of the antigen-antibody complex of influenza virus neuraminidase heads and NC10 MFab (Tulloch et al., 1986, J.Mol. Biol. 190, 215-225; Malby et al., 1994, Structure, 2, 733-746).

#### Image processing

Electron micrographs showing particles in a limited number of identifiable projections were chosen for digitisation. Micrographs were digitised on a Perkin-Elmer model 1010 GMS PDS flatbed scanning microdensitometer with a scanning aperture (square) size of 20 mm and

stepping increment of 20 mm corresponding to a distance of 0.2 nm on the specimen. Particles were selected from the digitised micrograph using the interactive windowing facility of the SPIDER image processing system (Frank et al., 1996, *J. Struct. Biol.* 116, 190-199). Particles were scaled to an optical density range of 0.0 - 2.0 and aligned by the PSPC reference-free alignment algorithm (Marco et al., 1996, *Ultramicroscopy*, 66, 5-10). Averages were then calculated over a subset of correctly aligned particles chosen interactively as being representative of a single view of the particle. The final average image presented here is derived from a library of 94 images.

#### Biochemical characterization of expressed hIR -11 ectodomain

The recombinant protein examined corresponded to the the first 914 residues of the 917 residue ectodomain of the exon -11 form of the human insulin receptor (Ullrich et al., 1986, Nature 313, 756-761). Expressed protein was shown, by SDS-PAGE and autoradiography of immunoprecipitated product from metabolically labelled cells, to exist as a homodimeric complex of  $\sim$ 270 - 320 kDa apparent mass, which dissociated under reducing conditions into monomeric  $\alpha$  and  $\beta$ ' subunits of respective apparent mass  $\sim$ 120 kDa and  $\sim$ 35 kDa (data not shown).

Purified hIR -11 ectodomain, expressed in Lec8 cells and purified by affinity chromatography on an insulin affinity column, ran as a symmetrical peak on a Superdex S200 gel filtration column (Figure 16). The protein eluted with an apparent mass of ~400 kDa, calculated from a standard curve generated by the elution positions of standard proteins (not shown). As expected for protein expressed in Lec 8 cells, whose glycosylation defect produces truncated oligosaccharides (Stanley, 1989, . Molec. Cellul. Biol. 9, 377-383), this value is less than the apparent mass (450 - 500 kDa) reported for hIR +11 ectodomain expressed in wild-type CHO-K1 cells (Johnson et al., 1988, Proc. Natl Acad. Sci USA 85, 7516-7520; Cosgrove et al., 1995, Protein Expression and Purification 6, 789-798).

Radioassay of insulin binding to purified ectodomain gave linear Scatchard plots and Kd values of 1.5 - 1.8 x 10-9 M, similar to the values of 2.4 - 5.0 x 10-9 M reported for the hIR -11 ectodomain (Andersen et al., 1990, Biochemistry 29, 7363-7366; Markussen et al., 1991, J. Biol. Chem. 266, 18814-18818; Schaffer, 1994, Eur. J. Biochem. 221, 1127-1132) and the values of ~1.0 - 5.0 x 10-9 M reported for the hIR +11 ectodomain (Schaefer et al., 1992, J. Biol. Chem. 267, 23393-23402; Whittaker et al., 1994, Molec.

Endocrinol. 8, 1521-1527; Cosgrove et al., 1995, Protein Expression and Purification 6, 789-798).

#### Expression of hIGF-1R ectodomain

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Cloning, expression and purification of this protein used elements common to those described for hIR -11 ectodomain (Cosgrove et al., 1995, Protein Expression and Purification 6, 789-798) and resulted in purified product that was recognised by receptor-specific Mabs 17-69, 24-31 and 24-60 (Soos et al., 1992, J. Biol. Chem. 267, 12955-63) and was composed of α and β' subunits of mass similar to those of hIR ectodomain (unpublished data). Preparation of hIR -11 ectodomain/MFab complexes

A complex of hIR -11 ectodomain and Fab from antibody 83-14 eluted as a symmetrical peak of 460 -500 kDa (Figure 16), as did complexes generated from a mixture of hIR -11 ectodomain with Fab from antibody 18-44 and a mixture of hIR -11 ectodomain with Fab 83-7 (not shown). A cocomplex of ectodomain with Fabs from antibodies 18-44 and 83-14 eluted at 620 kDa (Figure 12), as did a co-complex with MFabs 83-14/83-7 and another with MFabs 83-7/18-44 (not shown). A complex of hIR -11 ectodomain with all three MFab derivatives, 18-44, 83-7 and 83-14, eluted at an apparent mass of ~ 710 kDa (Figure 16).

#### **Electron microscopy**

#### Imaging of hIR -11 and hIGF-1R ectodomains

Single-molecule imaging of undecorated dimeric hIR -11 ectodomain was carried out under a variety of negative staining conditions, which emphasised different aspects of the structure of the molecular envelope. The least aggressive or penetrative stain was potassium phosphotungstate (KPT), which revealed consistent globular particles with very little internal structure other than a suggestion of a division into two parallel bars. Staining with methylamine tungstate also revealed the parallel bar images, as shown in Figure 17a.

Further investigation using progressively more penetrative, but also potentially more disruptive, stains confirmed the observations above. Staining with uranyl acetate and uranyl formate showed the separation of the parallel bars most clearly (Figure 17b), but uranyl acetate showed evidence of disrupting the structure of the particles, i.e. a decrease in the consistency of the particle shape and a tendency for particles to look unravelled or denatured despite having been subjected to chemical cross-linking prior to

staining. In areas of thicker stain, parallel bars predominated (Figure 17b), whereas in more thinly stained regions, U-shaped particles could be identified, sometimes outnumbering the parallel-bar structures (Figure 18a). An averaged image of the parallel bars seen by staining hIR -11 ectodomain with uranyl formate is shown as an insert in Figure 17b.

In Figures 17c and 18b, images of hIGF-1R ectodomain are shown for comparison with Figure 17b and 18a, respectively, under similar staining conditions.

#### Imaging of hIR -11 ectodomain complexed with 83-7 MFab

This complex was particularly noteworthy for the consistency of the form of the particles, especially under the gentler staining conditions afforded by stains such as KPT and methylamine tungstate. The particles were interpreted as having been restricted in the views they presented, after air-drying on the carbon support film, by the almost diametrically opposite binding of the two Fab arms to the antigen to form a highly elongated complex structure. Under these conditions three distinct views could be recognised as shown in Figure 19. Two views (interpreted as top-down/bottom-up) show the Fab arms displaced clockwise or anti-clockwise as extensions of the parallel plates with two-fold symmetry. The third view shows an image with the two Fab arms in line roughly through the centre of the receptor on its opposite sides, interpreted as a side projection of binding half-way up the plates (Figure 19).

Figure 20 shows a field of particles of hIR -11 ectodomain complexed with 83-7 MFab, stained with uranyl formate. The use of the more aggressive uranyl stains operating at lower pHs revealed internal structure of the molecular envelope at the expense of consistency of the particle morphology. For example, staining with uranyl acetate or uranyl formate showed that parallel bars can be seen in particles in which the Fab arms are displaced either clockwise or anticlockwise but not where the intermediate central or axial position of the two Fab arms is presented in projection. These observations show 83-7 MFab binding roughly half-way up the side-edge of each hIR -11 ectodomain plate. The epitope recognised by Mab 83-7 has been mapped to the cys-rich region, residues 191-297, by analysis of chimeric receptors (Zhang and Roth, 1991, *Proc. Natl. Acad. Sci. USA* 88, 9858-9862).

# Imaging of hIR -11 ectodomain complexed with either 83-14 MFab or 18-44 MFab

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Figure 21a shows the complexes formed with Fabs from the most insulin-mimetic antibody Mab 83-14. Projections showing the Fab arms bound to and extending out from near the base of the U-shaped particles can be identified. A second field of particles (Figure 21b) shows objects composed of two parallel bars as observed for the undecorated ectodomain, with Fab arms projecting obliquely from diametrically opposite extremities. Similar but less definitive images were also seen when MFab 18-44 was bound to hIR -11 ectodomain (not shown). The epitope for Mab 83-14 is between residues 469-592 (Prigent et al., 1990) in the connecting domain. This domain contains one of the disulphide bonds (Cys524-Cys524) between the two monomers in the IR dimer (Schaffer and Ljungqvist, 1992, Biochem. Biophys. Res. Commun. 189, 650-653). The epitope for Mab 18-44 is a linear epitope, residues 765-770 (Prigent et al., 1990, . J. Biol. Chem. 265, 9970-9977) in the  $\beta$ -chain, near the end of the insert domain (O'Bryan et al., 1991, Mol. Cell. Biol. 11, 5016-5031). The insert domain contains the second disulphide bond connecting the two monomers in the IR dimer (Sparrow et al., 1997, J. Biol. Chem., 272, 29460-29467).

## Imaging of hIR -11 ectodomain co-complexed with two different MFabs per monomer

The double complex of hIR -11 ectodomain with MFabs 83-7 and 18-44 was stained with 2% KPT at pH 6.0, and revealed the molecular envelopes shown in Figure 22. The particle appears complex in shape and can assume a number of different orientations on the carbon support film, giving rise to a number of different projections in the micrograph. The predominant view is of an asymmetric X-shape (some examples circled). It shows the 83-7 MFab arms bound at opposite ends of the parallel bars with the two 18-44 MFabs appearing as shorter projections extending out from either side of each ectodomain.

Images of the double complex of hIR -11 ectodomain with 83-7 and 83-14 MFabs gave X-shaped images similar to those seen with the 83-7/18-44 double complex (not shown). In contrast the double complex of hIR -11 ectodomain with 18-44 and 83-14 MFabs did not present the characteristic asymmetric X-shapes described above (images not shown). Instead, the molecular envelope appeared to be elongated in many views, with only an

occasional X-shaped projection. While a detailed interpretation of these images would be premature, it is clear that MFabs 18-44 and 83-14, two of the more potent insulin mimetic antibodies (Prigent et al., 1990, *J. Biol.* 

Chem. 265, 9970-9977), can bind simultaneously to the receptor.

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# Imaging of hIR -11 ectodomain co-complexed with three different MFabs per monomer

Figure 23 shows a field of particles from a micrograph of hIR -11 ectodomain complexed simultaneously with MFabs 83-7, 83-14 and 18-44. In the thicker stain regions the molecular envelope is X-shaped, and looks very similar to that of the double complexes of hIR -11 ectodomain with either 83-7 and 18-44 or 83-7 and 83-14. However, in the more thinly stained regions, particles of greater complexity are visible and it is possible occasionally to identify that there are in fact more than four MFabs bound to the ectodomain dimer.

The single-molecule imaging of hIR -11 ectodomain presented here suggests a molecular envelope for this dimeric species significantly different from that of any previously published study. However, an unequivocal determination of the molecular envelope even from the present study is not entirely straightforward. A major complicating factor here has been the relative fragility of the expressed ectodomain when exposed to the rigors of electron microscope preparation by negative staining. For example, staining with potassium phosphotungstate (KPT, pH 6.0-7.0) frequently suggested a denaturation of the dimeric molecules, but when appropriate conditions were satisfied, good seemingly interpretable molecular envelope images were achieved; staining with methylamine tungstate (pH  $\sim$  7.0) supported the best KPT molecular envelope images, but had the suggestion of a swelling of the molecular structure at neutral pH; and the acid-pH stains of uranyl acetate ( pH  $\sim$ 4.2) and uranyl formate (pH $\sim$ 3.0), with their ability to penetrate the ectodomain structure, appeared to illuminate not so much the molecular envelope as the zones of high projected protein density within the dimer.

An amalgam of impressions from these various staining regimens has led to the following interpretation of single-molecule images of these undecorated, or naked, dimers: the predominant dimeric molecular image encountered here has been that of 'parallel bars' of projected protein density. This view is so predominant, indeed, that it suggests there is either a single preferred orientation of the molecules on the glow-discharged carbon support

film, or that this impression of parallel bars of density may represent a mixture of superficially similar structure projections, with the subtleties of these different projections being masked by the relatively coarse resolution of this single-molecule direct imaging. The impression of parallel bars of projected protein density is particularly predominant in regions of thicker negative stain. A second view of the molecular envelope, appreciably less well represented in regions of thicker stain but predominant in regions of thin staining, is that of 'open' U's, or V's. These two views of hIR -11 ectodomain were supported by the single-molecule imaging of hIGF-1R ectodomain under comparable conditions of negative staining.

If the assumption is made that these two recognisable projected views, that of parallel bars and of open U's/V's, are different views of the same dimeric molecule, an assumption strongly supported by the MFab complex imaging, a coarse model of the molecular envelope can be rationalized as in the schematic Figure 24. The model structure is roughly that of a cube, composed of two almost-parallel plates of high protein density, separated by a deep cleft of low protein main-chain and side-chain density able to be penetrated by stain, and connected by intermediate stain-excluding density near what is assumed here to be their base ( that is, nearest the membrane-anchoring region). The width of the low-density cleft appears to be of the order of 30-35Å, sufficient to accommodate the binding of the insulin molecule of diameter ca. 30Å, although we have no electron microscopical evidence to support insulin-binding in this cleft at this stage.

It has been established through imaging of bound 83-7 MFab that there is a dimeric two-fold axis normal to the membrane surface between these plates of density. Occasionally, dimer images display a relative displacement of the bars of density, interpreted here as a limited capacity for a shearing of the interconnecting zone between the two plates along their horizontal axis parallel to the membrane; other images show bars skewed from parallel, implying a limited capacity for the plates to rotate independently around the two-fold axis, again via this interconnecting zone. These two observations each suggest a relatively flexible connectivity between the dimer plates in the membrane-proximal region of intermediate protein density, which could possibly contribute to the transmembrane signalling process.

The approximate overall measured dimensions of the ectodomain dimer depicted in Figure 24 are 110 x 90 x 120Å, calibrated against the dimensions of imaged influenza neuraminidase heads, known from the solved X-ray structure (Varghese et al., 1983, Nature 303, 35-40). It can be noted that there is a compatibility here between the molecular weights and molecular dimensions of these two molecular species: the compact tetrameric influenza neuraminidase heads of Mr ~200 kDa occupy a volume almost 100 x 100 x 60 Å; the more open dimeric insulin receptor ectodomains of similar Mr ~240 kDa imaged here occupy a volume approximately 110 x 90 x 120 Å , roughly twice that of the neuraminidase heads, accommodating the slightly higher molecular weight and substantial central low-density cleft.

The low-resolution roughly cubic compact structure proposed here differs substantially from the T-shaped model proposed by Christiansen et al. (1991, Proc. Natl. Acad. Sci. U. S. A. 88, 249-252) and Tranum-Jensen et al., (1994, J. Membrane Biol. 140, 215-223) for the whole receptor and the elongated model proposed by Schaefer et al. (1992, J. Biol. Chem. 267, 23393-23402) for soluble ectodomain. Significantly, those previous studies did not provide any convincing independent electron microscopical evidence that their imaged objects were in fact insulin receptor.

In the present study, the identity of the imaged molecules as hIR -11 ectodomain has been confirmed by imaging complexes of the dimer with Fabs of the three well-established conformational Mabs against native hIR, 83-7, 83-14 and 18-44 (Soos et al.,1986, Biochem. J. 235, 199-208; 1989, Proc. Natl Acad. Sci. USA 86, 5217-5221), bound singly and in combination. In all these instances, virtually every particle in the field of view exhibited MFab decoration through binding to conformational epitopes, establishing not only the identity of the imaged particles but also the conformational integrity of the expressed ectodomains. Furthermore, the cleanliness and uniformity of these hIR -11 ectodomain preparations, both naked and decorated, visualised here by electron microscopy demonstrate their high suitability for X-ray crystallization trials.

The known flexibility of the Fab arms exacerbates image-to-image variability beyond the limited extent already described for the undecorated dimeric ectodomains, complicating any precise interpretation of these antigen-antibody complexes. Such molecular flexibility also renders largely impractical any single-molecule computer image averaging to facilitate image

interpretation, progressively more so with the higher order antigen-antibody complexes studied here.

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The most readily interpretable of these images, showing least imageto-image variability, are those of 83-7 MFab bound to dimers where, fortuitously, the antigen-antibody complex is constrained in its degrees of rotational freedom on the carbon support film. Many projected images show the two Fab arms in line roughly through the centre of the antigen on its opposite sides (Figure 19, arrowed examples), interpreted as a side projection of binding half-way up the plates from their membrane-proximal base. Other sub-sets of images (Figure 19, circled examples ) show the two Fab arms still parallel but displaced clockwise or anticlockwise with 2-fold symmetry, each Fab approximating an extension of one of the parallel bars of antigen density, interpreted here as representing top or bottom projections along the 2-fold axis. The third projection, along the axis of the Fab arms, could not be sampled here because of the constraining geometry of this molecular complex. These observations suggest binding of 83-7 MFab roughly half-way up the side-edge of the hIR -11 ectodomain plate. This then allows an initial attempt at spatially mapping the 83-7 MFab epitope, which has been sequence-mapped to residues 191-297 in the cys-rich region of the insulin receptor (Zhang and Roth, 1991, Proc. Natl. Acad. Sci. USA 88, 9858-9862). The spatial separation and relative orientations of the two binding epitopes of Mab 83-7 on the hIR -11 ectodomain dimer as indicated here appear inconsistent with the proposal that Mab 83-7 could bind intramolecularly to hIR (O'Brien et al., 1987, Biochem J. 6, 4003-4010).

Decoration of the ectodomain dimer with 83-7 MFab established that the two plates of high protein-density are arranged with 2-fold symmetry. Decoration with either 83-14 or 18-44 MFab, on the other hand, allowed sampling of the third projection of the ectodomain dimer precluded by 83-7 MFab binding. Significantly, this third view established unequivocally the U-shaped projection of the hIR -11 ectodomain dimer, something which was only able to be assumed with the undecorated ectodomain images. Further, this projection has allowed a rough spatial mapping close to the base of the U-shaped dimer for the epitopes recognised by 83-14 MFab (residues 469-592, connecting domain) and 18-44 MFab (residues 765-770, b-chain insert domain; exon 11 plus numbering, Prigent et al., 1990, J. Biol. Chem. 265, 9970-9977).

Inherent in the model structure presented in Figure 20 is the implication that, with the two-fold axis aligned normal to the membrane surface, the mouth of the low-density cleft where insulin binding may occur would lie most distant from the transmembrane anchor, whilst the zone of intermediate density connecting the two high-density plates would be in close proximity to the membrane. It follows, in this model, that the L1/cysrich/L2 domains(Bajaj et al., 1997, Biochim. Biophys. Acta 916, 220-226; Ward et al.,1995, Proteins: Struct., Funct., Genet. 22, 141-153), which comprise much of the insulin-binding region (see Mynarcik et al., 1997, . J. Biol. Chem. 272, 2077-2081), most probably lie in the membrane-distal upper halves of the two plates, whilst the membrane-proximal lower halves contain the connecting domains, the fibronectin-type domains, the insert domains and the interchain disulphide bonds (Schaffer and Ljungqvist, 1992, Biochem. Biophys. Res. Commun. 189, 650-653; Sparrow et al., 1997, J. Biol. Chem., 272, 29460-29467). Such a disposition of domains is supported by the images seen with the single MFab decoration, the 83-7 MFab epitope in the cys-rich region being spatially mapped roughly half-way up the side-edge of the ectodomain plates, and the 83-14 and 18-44 MFab epitopes (connecting domain and \beta-chain insert domain, respectively) being mapped near the base of the plates. Our preference is for a single a-b¢ monomer to occupy a single plate, although the possibility of a single monomer straddling the two plates of protein density cannot be discounted.

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The more complex images involving co-binding of two, and even more so of all three, MFabs to each monomer of the ectodomain dimer (Figures 22 and 23) are not easily interpretable with respect to relative domain arrangements within the monomer at present, not least of all because of the difficulty of finding conditions of negative staining that will simultaneously maintain the integrity of the Fab binding while highlighting recognisable and reproducible details of the internal structure of the dimeric IR ectodomain.

The data presented here demonstrate the ability of single-molecule imaging to give an initial insight into the topology of multidomain structures such as the ectodomain of hIR, and the value of combining this technique with that of either single or multiple monoclonal Fab attachment per monomer as a potential means of epitope (and domain ) mapping of the structure. By imaging Fab complexes of other members of the family (such as

hIGF-1R ectodomain) and combining available sequence-mapped epitope information with that presented here, a more comprehensive understanding of domain arrangements within the IR family ectodomains should be forthcoming.

It will be appreciated by persons skilled in the art that numerous variations and/or modifications may be made to the invention as shown in the specific embodiments without departing from the spirit or scope of the invention as broadly described. The present embodiments are, therefore, to be considered in all respects as illustrative and not restrictive

Dated this twenty-seventh day of November 1997

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COMMONWEALTH SCIENTIFIC
AND INDUSTRIAL RESEARCH
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Patent Attorneys for the Applicant:

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251       13.277       61.488         298       14.074       60.248         713       11.849       61.237         718       11.005       60.485         5561       15.328       62.821         357       15.467       62.933         392       16.356       62.950         869       17.691       63.196         919       18.587       63.813         365       19.807       64.466         459       20.683       54.959         340       20.970       64.141         399       18.320       62.896         639       19.224       61.354         251       17.845       61.405         688       18.356       60.174         585       17.409       59.787         502       16.983       60.610         531       17.029       58.531         486       16.119       58.127         586       16.818       57.134         287       16.881       57.134         286       17.936       53.815         451       18.921       54.366         522       17.936       53.81	8.785       53.34         2.939       9.448       52.33         3.625       10.841       52.65         10.930       51.19         10.930       51.19         1403       12.789       52.41         12.789       52.41         1606       9.077       57.71         1441       7.138       57.53         1685       7.209       58.78         1685       7.209       58.78         149       8.824       58.16         149       8.824       58.16         149       8.824       58.16         149       8.824       58.16         120       4.492       59.58         120       4.492       59.58         120       4.492       59.58         120       4.492       59.58         1372       2.483       58.33         409       5.413       61.95         3.72       2.483       58.33         409       5.413       61.95         3.567       58.67       58.29         3.372       2.483       58.33         409       5.413       61.95
1.00 26.05 1.00 22.27 1.00 21.10 1.00 26.19 1.00 23.67 1.00 23.69 1.00 19.28 1.00 17.36 1.00 20.66  1.00 26.10 1.00 25.08 1.00 20.44 1.00 20.35 1.00 20.44 1.00 19.17 1.00 20.35 1.00 19.17 1.00 20.35 1.00 19.17 1.00 21.44 1.00 21.42 1.00 20.35 1.00 19.17 1.00 22.05 1.00 21.44 1.00 21.42 1.00 20.45 1.00 21.44 1.00 21.42 1.00 20.43 1.00 19.50 1.00 19.50 1.00 19.50 1.00 19.50 1.00 19.50 1.00 27.73 1.00 29.55 1.00 27.42 1.00 20.63 1.00 19.50 1.00 27.75 1.00 27.42 1.00 20.63 1.00 19.50 1.00 27.77 1.00 29.55 1.00 27.42 1.00 20.63 1.00 19.54 1.00 20.63 1.00 19.54 1.00 27.77 1.00 29.55 1.00 27.42 1.00 20.63 1.00 19.54 1.00 21.37 1.00 29.55 1.00 27.42 1.00 20.63 1.00 19.54 1.00 21.37 1.00 23.29 1.00 20.63 1.00 37.44 1.00 23.29 1.00 23.58 1.00 36.56 1.00 37.08 1.00 37.08 1.00 37.44 1.00 38.01 1.00 39.45 1.00 37.44 1.00 37.46	1.00 45.71 1.00 46.54 1.00 44.56 1.00 38.05 1.00 34.60 1.00 37.23 1.00 34.22 1.00 32.23 1.00 27.55 1.00 29.77 1.00 35.63 1.00 26.22 1.00 34.55 1.00 26.83 1.00 26.83 1.00 27.88 1.00 17.88 1.00 10.19
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Figura 1

MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM
405 CE2 TYR 406 CZ TYR 407 OH TYR 409 C TYR 410 O TYR 411 N ARG 413 CA ARG 414 CB ARG 415 CG ARG 416 CD ARG 417 NE ARG 420 NH1 ARG 423 NH2 ARG 426 C ARG 427 O ARG 428 N PHE 430 CA PHE 431 CB PHE 432 CG PHE 433 CD1 PHE 434 CD2 PHE 435 CE1 PHE 436 CE2 PHE 437 CZ PHE 438 C PHE 439 O PHO 441 CD PRO 442 CA PRO 441 CD PRO 442 CA PRO 444 CG PRO 444 CG PRO 445 C PRO 445 C PRO 446 O PRO 447 N LYS 458 C LYS 459 O LYS 450 CB LYS 451 CG LYS 451 CG LYS 452 CD LYS 453 CE LYS 454 NZ LYS 456 CD1 LEU 466 CD2 LEU 467 C LEU 467 C LEU 468 C THR 471 CA THR 472 CB THR 473 CG2 THR 475 CG2 THR 476 C THR 477 O THR 478 N VAL 480 CA VAL 481 CB VAL 482 CG1 VAL 483 CG2 VAL 484 C VAL 485 O LIE	125 CG2 ILE 326 CG1 ILE 327 CD1 ILE 328 C ILE 329 O ILE 330 N SER 3312 CA SER 3314 OG SER 3316 C SER 3317 O SER 3316 CA LYS 3318 N LYS 3340 CA LYS 341 CB LYS 341 CB LYS 342 C LYS 341 CB ALA 341 CB ALA 341 CB ALA 342 C GLU 353 CB GLU 354 CG GLU 355 C GLU 355 CB GLU 356 C GLU 357 O GLU 358 N ASP 360 CA ASP 361 CB ASP 361 CB ASP 362 CC ASP 363 OD1 ASP 364 OD2 ASP 365 C ASP 366 O ASP 367 N TYR 369 CA TYR 370 CB TYR 371 CG TYR 371 CG TYR 372 CD1 TYR 373 CE1 TYR 374 CD2 TYR 375 CE2 TYR 376 CZ TYR 377 OH TYR 379 C TYR 370 C TYR 371 C TYR
4333344444444455555555555556666667777777777	344445555555666667777788888889999999999999999
51.438       -0.818         52.242       -0.731         48.181       -2.148         47.054       -2.639         48.728       -1.446         47.998       -1.192         47.794       -2.464         47.337       -2.191         47.561       -3.382         46.299       -3.944         45.831       -3.778         46.532       -3.069         44.663       -4.312         48.794       -0.223         49.907       -0.544         48.871       2.011         48.865       3.286         49.077       3.019         50.242       3.389         48.121       2.354         50.459       3.101         48.337       2.066         49.508       2.444         48.254       2.222         47.462       3.113         48.638       1.381         49.619       0.289         48.114       1.493         48.651       2.759         48.751       2.759         48.651       2.759         48.749       4.986         52.617	45.030
53.2421 53.	48.5768 46.2005 46.2005 46.2005 46.2005 46.2005 46.2005 47.368 47.368 47.47.4918 47.47.4918 47.47.4918 47.47.4918 47.47.4918 47.48.5798 47.48.5798 47.48.5798 47.4918 47.4918 47.49.612 47.49
1.00 49.57 1.00 50.74 1.00 52.29 1.00 49.86 1.00 48.04 1.00 45.99 1.00 43.34 1.00 50.00 1.00 54.95 1.00 58.18 1.00 60.47 1.00 61.57 1.00 64.67 1.00 41.23 1.00 41.95 1.00 36.81 1.00 33.21 1.00 28.46 1.00 32.20 1.00 32.67 1.00 31.49 1.00 31.66 1.00 31.15 1.00 32.77 1.00 34.39 1.00 36.69 1.00 35.29 1.00 36.69 1.00 35.53 1.00 36.69 1.00 35.53 1.00 40.65 1.00 40.82 1.00 40.82 1.00 40.82 1.00 40.82 1.00 40.82 1.00 40.82 1.00 40.82 1.00 40.82 1.00 40.83 1.00 26.62 1.00 29.34 1.00 33.36 1.00 26.62 1.00 29.34 1.00 33.36 1.00 26.52 1.00 29.34 1.00 33.36 1.00 26.52 1.00 29.34 1.00 33.36 1.00 26.52 1.00 29.34	1.00 89.97 1.00 85.71
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MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM
563 CE2 PHE 564 CZ PHE 565 C PHE 565 C PHE 566 O PHE 567 N ARG 570 CB ARG 571 CG ARG 571 CG ARG 572 CD ARG 573 NE ARG 575 CZ ARG 576 NH1 ARG 579 NH2 ARG 582 C ARG 583 O ARG 584 N VAL 586 CA VAL 587 CB VAL 587 CB VAL 588 CG1 VAL 589 CG2 VAL 591 O ALA 592 N ALA 595 C ALEU 591 O C CA 601 C GLY 602 O GLY 603 N LEU 605 CA LEU 606 CB LEU 607 CG LEU 608 CD1 LEU 610 C LEU 611 O LEU 611 O LEU 612 N GLU 615 CG GLU 616 CB GLU 617 CD GLU 618 CE GLU 619 OE2 GLU 610 C SER 626 OG SER 627 CB SER 628 C SER 628 C SER 629 O SER 629 O SER 629 C SER 620 CB LEU 631 CB LEU 633 CB LEU 633 CB LEU 634 CG CB 635 CD1 LEU 636 CD2 LEU 637 C GLY 638 CD1 LEU 639 CB SER 629 O SER 629 O SER 629 O SER 629 C SER 629 C SER 620 CB CD2 LEU 631 CB LEU 633 CB LEU 634 CG GLY 635 CD1 LEU 636 CD2 LEU 637 C GLY 638 CD1 LEU 639 CB SER 629 C	*** *** *** *** *** *** *** *** *** **
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35.670 37.375 38.563 36.436 36.761 35.973 35.739 36.036 36.603 35.887 34.560 36.508 37.993 39.054 37.858 38.973 39.571 40.461 40.373	42.410 42.347 42.914 42.704 41.547 43.450 44.595 44.506 41.6405 42.4205 42.4205 43
10.640 7.251 6.497 6.398 6.398 6.875 6.975 6.875 6	12.877 13.595 11.441 10.630 14.924 14.938 16.022 17.284 18.250 18.826 17.554 18.002 19.180 17.310 17.310 17.310 17.310 17.310 17.310 17.310 17.310 17.310 17.310 17.310 17.310 17.310 17.310 18.409 19.829 19.724 11.581 16.407 16.455 16.407 16.455 16.407 16.455 16.407 16.455 16.407 16.455 16.407 16.455 17.888 15.979 17.819 18.212 17.886 15.979 17.819 18.212 17.885 18.325 11.428 11.934
5.1489 5.1489 7.6.859 7.6.8	1.865 1.893 1.993
1.00 29.70 1.00 30.40 1.00 34.00 1.00 39.70 1.00 32.00 1.00 29.35 1.00 28.94 1.00 41.50 1.00 53.38 1.00 56.20 1.00 54.54 1.00 59.91 1.00 29.92 1.00 34.29 1.00 27.82 1.00 27.82 1.00 27.82 1.00 27.82 1.00 27.82 1.00 27.82 1.00 27.82 1.00 27.82 1.00 34.37 1.00 30.46 1.00 34.37 1.00 31.76	1.00 18.06 1.00 17.87 1.00 20.56 1.00 19.01 1.00 12.40 1.00 18.06 1.00 18.83 1.00 18.83 1.00 16.95 1.00 20.53 1.00 17.67 1.00 25.70 1.00 25.70 1.00 25.70 1.00 28.12 1.00 28.12 1.00 23.84 1.00 28.12 1.00 26.32 1.00 23.93 1.00 23.46 1.00 23.93 1.00 23.46 1.00 22.72 1.00 22.72 1.00 22.72 1.00 22.77 1.00 27.17 1.00 21.86 1.00 22.82 1.00 22.77 1.00 21.86 1.00 16.41 1.00 16.41 1.00 16.41 1.00 16.41 1.00 16.41 1.00 16.41 1.00 22.77 1.00 27.77 1.00 27.77 1.00 27.77 1.00 27.77 1.00 23.36 1.00 24.00 1.00 23.70 1.00 24.00 1.00 23.70 1.00 23.70 1.00 24.00 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 23.70 1.00 24.05 1.00 25.00 1.00 23.70
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ATOM 644 N ASP 68 ATOM 646 CA ASP 68 ATOM 650 CB ASP 68 ATOM 650 OD2 ASP 68 ATOM 650 OD2 ASP 68 ATOM 651 C ASP 68 ATOM 652 O ASP 68 ATOM 655 CA LEU 69 ATOM 655 CA LEU 69 ATOM 655 CB LEU 69 ATOM 656 CB LEU 69 ATOM 657 CG LEU 69 ATOM 658 CD1 LEU 69 ATOM 660 C LEU 69 ATOM 660 C LEU 69 ATOM 666 CB PHE 70 ATOM 667 CD1 PHE 70 ATOM 666 CB PHE 70 ATOM 667 CD1 PHE 70 ATOM 667 CD1 PHE 70 ATOM 667 CD1 PHE 70 ATOM 667 CD2 PHE 70 ATOM 667 CD1 PHE 70 ATOM 670 CE2 PHE 70 ATOM 671 CZ PHE 70 ATOM 673 O PHE 70 ATOM 675 CD PRO 71 ATOM 676 CA PRO 71 ATOM 676 CA PRO 71 ATOM 677 CB PRO 71 ATOM 679 C PRO 71 ATOM 680 O PRO 71 ATOM 681 N ASN 72 ATOM 683 CA ASN 72 ATOM 684 CB ASN 72 ATOM 685 CG ASN 72 ATOM 687 CD1 LEU 73 ATOM 689 C ASN 72 ATOM 699 C LEU 73 ATOM 700 O LEU 73 ATOM 701 N THR 74 ATOM 703 CA THR 74 ATOM 709 C THR 74 ATOM 710 N VAL 75 ATOM 715 CG2 VAL 75 ATOM 716 C VAL 75 ATOM 717 C VAL 75 ATOM 718 N ILE 76 ATOM 719 C THR 74 ATOM 719 C THR 74 ATOM 710 N VAL 75 ATOM 716 C VAL 75 ATOM 717 C VAL 75 ATOM 718 N ILE 76 ATOM 718 N ILE 76 ATOM 719 C ALL 75 ATOM 716 C VAL 75 ATOM 717 C VAL 75 ATOM 718 N ILE 76 ATOM 718 N ILE 76	42.148 -3.116 60.675 1.00 39.06 41.803 -3.788 59.348 1.00 38.97 41.035 -5.095 59.528 1.00 43.61 40.891 -5.601 60.666 1.00 47.80	AAAA ATOM 803 CG TYR 83 AAAA ATOM 804 CD1 TYR 83 AAAA ATOM 805 CD2 TYR 83 AAAA ATOM 805 CD2 TYR 83 AAAA ATOM 806 C TYR 83 AAAA ATOM 806 C TYR 83 AAAA ATOM 807 O TYR 83 AAAA ATOM 808 N ASN 84 AAAA ATOM 810 CA ASN 84 AAAA ATOM 811 CB ASN 84 AAAA ATOM 812 CG ASN 84 AAAA ATOM 813 OD1 ASN 84 AAAA ATOM 813 OD1 ASN 84 AAAA ATOM 814 ND2 ASN 84 AAAA ATOM 817 C ASN 84 AAAA ATOM 818 O ASN 84 AAAA ATOM 819 N TYR 85 AAAA ATOM 819 N TYR 85 AAAA ATOM 821 CG TYR 85 AAAA ATOM 822 CB TYR 85 AAAA ATOM 822 CB TYR 85 AAAA ATOM 822 CC TYR 85 AAAA ATOM 825 CC1 TYR 85 AAAA ATOM 826 CD2 TYR 85 AAAA ATOM 827 CE2 TYR 85 AAAA ATOM 828 CZ TYR 85 AAAA ATOM 829 OH TYR 85 AAAA ATOM 820 CT TYR 85 AAAA ATOM 831 C TYR 85 AAAA ATOM 832 O TYR 85 AAAA ATOM 831 C TYR 85 AAAA ATOM 831 C TYR 85 AAAA ATOM 832 O TYR 85 AAAA ATOM 836 CB ALA 86 AAAA ATOM 837 C ALA 86 AAAA ATOM 838 O ALA 86 AAAA ATOM 836 CB ALA 86 AAAA ATOM 837 C ALA 86 AAAA ATOM 838 O ALA 86 AAAA ATOM 841 CA LEU 87 AAAA ATOM 842 CD LEU 87 AAAA ATOM 843 CG LEU 87 AAAA ATOM 844 CD1 LEU 87 AAAA ATOM 845 CD2 LEU 87 AAAA ATOM 846 C LEU 87 AAAA ATOM 846 C LEU 87 AAAA ATOM 847 O LEU 87 AAAA ATOM 848 N VAL 88 AAAA ATOM 848 N VAL 88 AAAA ATOM 846 CD LEU 87 AAAA ATOM 847 O LEU 87 AAAA ATOM 848 N VAL 88 AAAA ATOM 848 N VAL 88 AAAA ATOM 849 CD LEU 87 AAAA ATOM 840 CD LEU 87 AAAA ATOM 840 CD LEU 87 AAAA ATOM 846 CD LEU 87 AAAA ATOM 848 N VAL 88 AAAA ATOM 848 N VAL 88 AAAA ATOM 848	31.479 19.989 52.109 1.00 41.19 30.812 19.227 51.145 1.00 42.91 31.532 20.690 55.955 51.00 10.98 31.999 21.532 56.727 1.00 37.21 AAAA 30.655 19.811 56.399 1.00 26.48 AAAA 30.278 19.928 57.801 1.00 24.16 AAAA 28.218 21.108 56.880 1.00 35.15 AAAA 28.218 21.108 56.880 1.00 35.15 AAAA 28.049 22.325 57.058 1.00 35.15 AAAA 30.578 18.615 58.429 1.00 25.06 AAAA 30.11 18.309 55.511 1.00 25.06 AAAA 30.11 18.309 55.511 1.00 25.08 AAAA 31.371 17.820 57.740 1.00 25.57 AAAA 31.579 15.531 57.059 1.00 26.41 AAAA 31.579 15.531 57.059 1.00 32.86 AAAA 31.579 15.531 57.059 1.00 36.81 AAAA 30.211 15.601 56.429 1.00 35.46 AAAA 30.27.772 15.372 56.635 1.00 39.67 AAAA 31.371 17.820 57.740 1.00 38.56 AAAA 31.371 17.820 57.740 1.00 24.08 AAAA 31.579 15.531 57.059 1.00 36.81 AAAA 31.579 15.531 57.059 1.00 36.81 AAAA 31.579 15.532 57.059 1.00 37.07 AAAA 31.579 15.533 57.059 1.00 37.07 AAAA 31.371 17.820 60.58429 1.00 38.40 AAAA 31.371 17.820 15.4663 1.00 39.67 AAAA 31.371 17.820 15.4663 1.00 39.67 AAAA 31.371 17.820 15.4663 1.00 39.67 AAAA 31.373 10.617 6.605 1.00 37.07 AAAA 31.373 10.617 6.605 1.00 37.07 AAAA 31.3864 17.243 58.866 1.00 23.68 AAAA 31.319 15.387 59.784 1.00 19.80 AAAA 31.319 15.387 59.784 1.00 19.80 AAAA 31.403 15.169 60.544 1.00 18.47 AAAA 31.403 15.169 60.544 1.00 18.47 AAAA 31.403 15.169 59.252 1.00 18.47 AAAA 31.403 15.329 62.038 1.00 16.82 AAAA 31.424 11.569 59.252 1.00 17.46 AAAA 31.434 11.236 56.862 1.00 25.36 AAAA 31.434 11.236 56.862 1.00 25.36 AAAA 31.434 11.236 56.862 1.00 25.36 AAAA 31.345 8.293 61.318 1.00 15.41 AAAA 31.3450 10.935 55.579 1.00 27.52 AAAA 31.366 1.268 57.990 1.00 21.57 AAAA 31.368 1.268 57.990 1.00 21.57 AAAA 31.369 6.397 57.855 1.00 18.09 AAAA 31.373 10.617 60.359 1.00 15.94 AAAA 31.369 1.069 59.252 1.00 17.46 AAAA 31.369 1.069 59.252 1.00 17.46 AAAA 31.392 9.248 60.579 1.00 17.81 AAAA 31.392 9.248 60.579 1.00 17.81 AAAA 31.393 1.1667 60.564 1.00 18.09 AAAA 31.393 1.167 60.564 1.00 18.09 AAAA 31.399 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50 60.50
ATOM 721 CB ILE 76 ATOM 722 CG2 ILE 76 ATOM 723 CG1 ILE 76 ATOM 725 C ILE 76 ATOM 727 N ARG 77 ATOM 729 CA ARG 77 ATOM 729 CA ARG 77 ATOM 730 CB ARG 77 ATOM 731 CG ARG 77 ATOM 731 CG ARG 77 ATOM 732 CD ARG 77 ATOM 732 CD ARG 77 ATOM 732 CD ARG 77 ATOM 735 CZ ARG 77 ATOM 736 NH1 ARG 77 ATOM 736 NH1 ARG 77 ATOM 736 NH1 ARG 77 ATOM 736 NH2 ARG 77 ATOM 744 C C ARG 77 ATOM 743 O ARG 77 ATOM 744 N GLY 78 ATOM 748 O GLY 78 ATOM 748 O GLY 78 ATOM 749 N TRP 79 ATOM 751 CA TRP 79 ATOM 751 CA TRP 79 ATOM 752 CB TRP 79 ATOM 754 CD2 TRP 79 ATOM 756 CE3 TRP 79 ATOM 757 CD1 TRP 79 ATOM 758 NE1 TRP 79 ATOM 750 CZ TRP 79 ATOM 760 CZ2 TRP 79 ATOM 761 CZ3 TRP 79 ATOM 762 CH2 TRP 79 ATOM 763 C TRP 79 ATOM 764 N LYS ATOM 765 N LYS ATOM 765 N LYS ATOM 765 N LYS ATOM 767 CA LYS ATOM 768 CB LYS ATOM 769 CG LYS ATOM 770 CD LYS ATOM 771 CE LYS ATOM 771 CE LYS ATOM 772 N LYS ATOM 773 CD LYS ATOM 774 C LYS ATOM 775 CD LYS ATOM 775 CD LYS ATOM 776 C LYS ATOM 777 O LYS ATOM 778 C LYS ATOM 778 C LEU 81 ATOM 788 CB LEU 81 ATOM 789 CA LEU 81 ATOM 780 CA PHE 82 ATOM 790 CB PHE 8	37.832       12.671       64.548       1.00       15.15         37.815       13.532       63.303       1.00       12.00         38.462       11.314       64.195       1.00       14.57         38.165       14.720       66.028       1.00       22.48         36.979       14.951       65.953       1.00       22.55         39.019       15.613       66.478       1.00       20.68         38.570       16.912       66.923       1.00       23.08         39.786       17.718       67.429       1.00       23.55         40.503       17.036       68.605       1.00       22.68         41.648       17.842       69.102       1.00       24.68         41.223       19.034       69.825       1.00       11.24         40.728       19.027       71.066       1.00       32.80         40.587       17.881       71.725       1.00       31.08         40.436       20.178       71.681       1.00       36.79         37.717       17.784       66.005       1.00       23.75         36.959       18.358       63.881       1.00       24.91 <td< td=""><td>AAAA ATOM 880 CB GLU 91 AAAAA ATOM 881 CG GLU 91 AAAAA ATOM 883 CD GLU 91 AAAAA ATOM 883 CD GLU 91 AAAAA ATOM 883 CD GLU 91 AAAAA ATOM 885 C GLU 91 AAAAA ATOM 886 C GLU 91 AAAAA ATOM 886 C GLU 91 AAAAA ATOM 886 C GLU 91 AAAAA ATOM 887 N MET 92 AAAAA ATOM 889 CA MET 92 AAAAA ATOM 889 CA MET 92 AAAAA ATOM 890 CB MET 92 AAAAA ATOM 891 CG MET 92 AAAAA ATOM 892 SD MET 92 AAAAA ATOM 893 CE MET 92 AAAAA ATOM 893 CE MET 92 AAAAA ATOM 895 C MET 92 AAAAA ATOM 896 N THR 93 AAAAA ATOM 896 N THR 93 AAAAA ATOM 899 CB THR 93 AAAAA ATOM 899 CB THR 93 AAAAA ATOM 899 CB THR 93 AAAAA ATOM 900 CG THR 93 AAAAA ATOM 901 C THR 93 AAAAA ATOM 902 CG2 THR 93 AAAAA ATOM 903 C THR 93 AAAAA ATOM 904 C THR 93 AAAAA ATOM 905 N ASN 94 AAAAA ATOM 907 CA ASN 94 AAAAA ATOM 908 CB ASN 94 AAAAA ATOM 909 CG ASN 94 AAAAA ATOM 910 ND1 ASN 94 AAAAA ATOM 910 ND2 ASN 94 AAAAA ATOM 910 CD LYS 96 AAAAA ATOM 920 CG LEU 95 AAAAA ATOM 921 CD1 LEU 95 AAAAA ATOM 922 CD2 LEU 95 AAAAA ATOM 923 C LEU 95 AAAAA ATOM 924 C LEU 95 AAAAA ATOM 927 CA LYS 96 AAAAA ATOM 928 CB LYS 96 AAAAA ATOM 929 CG ASP 97 AAAAA ATOM 940 CA ASP 9</td><td>31.993</td></td<>	AAAA ATOM 880 CB GLU 91 AAAAA ATOM 881 CG GLU 91 AAAAA ATOM 883 CD GLU 91 AAAAA ATOM 883 CD GLU 91 AAAAA ATOM 883 CD GLU 91 AAAAA ATOM 885 C GLU 91 AAAAA ATOM 886 C GLU 91 AAAAA ATOM 886 C GLU 91 AAAAA ATOM 886 C GLU 91 AAAAA ATOM 887 N MET 92 AAAAA ATOM 889 CA MET 92 AAAAA ATOM 889 CA MET 92 AAAAA ATOM 890 CB MET 92 AAAAA ATOM 891 CG MET 92 AAAAA ATOM 892 SD MET 92 AAAAA ATOM 893 CE MET 92 AAAAA ATOM 893 CE MET 92 AAAAA ATOM 895 C MET 92 AAAAA ATOM 896 N THR 93 AAAAA ATOM 896 N THR 93 AAAAA ATOM 899 CB THR 93 AAAAA ATOM 899 CB THR 93 AAAAA ATOM 899 CB THR 93 AAAAA ATOM 900 CG THR 93 AAAAA ATOM 901 C THR 93 AAAAA ATOM 902 CG2 THR 93 AAAAA ATOM 903 C THR 93 AAAAA ATOM 904 C THR 93 AAAAA ATOM 905 N ASN 94 AAAAA ATOM 907 CA ASN 94 AAAAA ATOM 908 CB ASN 94 AAAAA ATOM 909 CG ASN 94 AAAAA ATOM 910 ND1 ASN 94 AAAAA ATOM 910 ND2 ASN 94 AAAAA ATOM 910 CD LYS 96 AAAAA ATOM 920 CG LEU 95 AAAAA ATOM 921 CD1 LEU 95 AAAAA ATOM 922 CD2 LEU 95 AAAAA ATOM 923 C LEU 95 AAAAA ATOM 924 C LEU 95 AAAAA ATOM 927 CA LYS 96 AAAAA ATOM 928 CB LYS 96 AAAAA ATOM 929 CG ASP 97 AAAAA ATOM 940 CA ASP 9	31.993

ATOM 973 CB TYR 101 ATOM 974 CG TYR 101 ATOM 975 CD1 TYR 101 ATOM 976 CE1 TYR 101 ATOM 977 CD2 TYR 101 ATOM 977 CD2 TYR 101 ATOM 979 CE2 TYR 101 ATOM 980 CH TYR 101 ATOM 980 CH TYR 101 ATOM 982 C TYR 101 ATOM 983 C TYR 101 ATOM 984 N ASN 102 ATOM 986 CA ASN 102 ATOM 988 CG ASN 102 ATOM 988 CG ASN 102 ATOM 989 CD1 ASN 102 ATOM 999 CD ASN 102 ATOM 999 CD ASN 102 ATOM 999 CD ASN 102 ATOM 999 CG LEU 103 ATOM 1000 CD1 LEU 103 ATOM 1001 CD2 LEU 103 ATOM 1004 N ARG 104 ATOM 1006 CA ARG 104 ATOM 1006 CA ARG 104 ATOM 1007 CB ARG 104 ATOM 1008 CG ARG 104 ATOM 1009 CD ARG 104 ATOM 1009 CD ARG 104 ATOM 1010 NE ARG 104 ATOM 1010 NE ARG 104 ATOM 1010 CB ASN 105 ATOM 1021 CG ASN 105 ATOM 1022 CB ASN 105 ATOM 1025 C ASN 105 ATOM 1029 CA ILE 106 ATOM 1029 CA ILE 106 ATOM 1033 CB ILE 106 ATOM 1034 C THR 107 ATOM 1035 CB THR 107 ATOM 1036 CB ARG 104 ATOM 1029 CA ILE 106 ATOM 1031 CG2 ILE 106 ATOM 1032 CB THR 107 ATOM 1034 C THR 107 ATOM 1035 C B TRR 107 ATOM 1036 CB ARG 104 ATOM 1037 CB TRR 107 ATOM 1038 CB THR 107 ATOM 1039 CB THR 107 ATOM 1034 C THR 107 ATOM 1035 C THR 107 ATOM 1036 CB ARG 108 ATOM 1037 CB ARG 108 ATOM 1038 CB ARG 108 ATOM 1039 CB THR 107 ATOM 1034 C THR 107	36.488 -1.023 68.722 1.00 29.53 37.890 -1.499 68.515 1.00 30.31 38.844 -1.386 69.525 1.00 29.96 40.133 -1.828 69.342 1.00 29.17 38.268 -2.065 67.306 1.00 30.96 30.543 3.510 67.100 1.00 30.96 40.480 -2.397 68.131 1.00 36.10 41.755 -2.908 67.934 1.00 44.43 36.834 1.319 69.470 1.00 32.61 36.387 1.213 70.628 1.00 35.65 37.804 2.173 69.146 1.00 34.59 38.368 3.060 70.156 1.00 32.17 39.883 2.919 70.153 1.00 34.62 40.355 1.842 71.107 1.00 38.21 40.312 2.018 72.328 1.00 39.16 40.783 0.709 70.560 1.00 37.20 37.928 4.506 69.909 1.00 33.315 38.322 5.443 70.621 1.00 34.93 37.089 4.681 68.889 1.00 33.71 35.879 5.969 67.157 1.00 27.71 35.499 7.287 66.474 1.00 24.01 36.520 8.393 66.654 1.00 14.91 35.342 6.955 65.028 1.00 26.24 35.598 6.465 69.623 1.00 30.84 34.479 5.966 69.711 1.00 30.84 34.479 5.966 69.711 1.00 30.84 36.027 7.422 70.437 1.00 28.81 35.135 7.877 71.502 1.00 26.99 35.954 7.732 72.815 1.00 31.56 36.358 6.291 73.131 1.00 30.84 36.577 9.269 71.408 1.00 26.71 33.618 9.562 72.120 1.00 26.99 35.425 4.508 74.671 1.00 28.81 35.964 1.737 77.552 1.00 26.99 35.425 4.508 74.671 1.00 40.69 34.835 12.187 69.278 1.00 22.794 35.169 7.626 77.1779 1.00 27.794 35.169 1.474 70.528 1.00 20.78 33.540 13.987 72.645 1.00 24.90 34.550 11.474 70.528 1.00 20.78 33.645 13.727 71.779 1.00 27.94 35.169 14.600 71.096 1.00 28.87 33.645 13.727 71.779 1.00 27.94 35.565 15.401 67.655 1.00 21.18 35.765 15.878 68.597 1.00 21.18 35.766 15.401 67.655 1.00 21.76 33.645 11.702 65.747 1.00 15.14 33.748 14.909 67.876 1.00 19.73 35.964 17.477 69.516 1.00 21.38 33.623 17.741 69.516 1.00 22.00 35.847 18.529 70.573 1.00 19.73 35.964 17.477 69.516 1.00 21.38 33.329 18.903 68.897 1.00 21.38 33.329 18.903 68.897 1.00 21.38 33.329 18.903 68.897 1.00 22.00	AAAA ATOM 1144 CA ALA 117 AAAA ATOM 1145 CB ALA 117 AAAA ATOM 1146 CB ALA 117 AAAA ATOM 1146 C ALA 117 AAAA ATOM 1147 O ALA 117 AAAA ATOM 1148 N ASP 118 AAAA ATOM 1148 N ASP 118 AAAA ATOM 1151 CB ASP 118 AAAA ATOM 1152 CG ASP 118 AAAA ATOM 1152 CG ASP 118 AAAA ATOM 1153 OD1 ASP 118 AAAA ATOM 1154 OD2 ASP 118 AAAA ATOM 1155 C ASP 118 AAAA ATOM 1156 O ASP 118 AAAA ATOM 1156 O ASP 118 AAAA ATOM 1157 N LEU 119 AAAA ATOM 1156 O ASP 118 AAAA ATOM 1156 O ASP 118 AAAA ATOM 1156 O ELEU 119 AAAA ATOM 1160 CB LEU 119 AAAA ATOM 1160 CB LEU 119 AAAA ATOM 1161 CG LEU 119 AAAA ATOM 1160 CB LEU 119 AAAA ATOM 1163 CD2 LEU 119 AAAA ATOM 1166 CD CYS 120 AAAA ATOM 1167 CD CYS 120 AAAA ATOM 1169 CC CYS 120 AAAA ATOM 1170 CC CC CYS 120 AAAA ATOM 1170 CC CC CYS 120 AAAA ATOM 1170 CC CC CC CYS 120 AAAA ATOM 1170 CC CC CC CYS 120 AAAA ATOM 1180 CC CC CC CYS 120 AAAA ATOM 1180 CC	27.796 -0.303 49.801 1.00 36.22 AA 27.0705 -1.512 49.636 1.00 36.19 AA 27.260 -2.093 48.290 1.00 38.96 AA 26.036 -3.065 51.118 1.00 41.77 AA 28.272 -3.003 51.106 1.00 39.97 AA 28.370 -4.080 52.108 1.06 38.19 AA 29.178 -5.158 51.719 1.00 44.53 AA 29.527 -5.314 50.222 1.00 52.11 AA 28.722 -6.072 49.628 1.00 56.23 AA 30.452 -4.683 49.645 1.00 55.85 AA 28.759 -3.647 53.471 1.00 33.86 AA 28.759 -3.647 53.471 1.00 33.85 AA 29.152 -4.487 54.258 1.00 33.86 AA 29.152 -4.487 54.258 1.00 33.86 AA 29.551 -3.33 56.478 1.00 27.83 AA 29.551 -0.332 56.478 1.00 22.15 AA 30.651 -0.392 57.220 1.00 22.15 AA 30.651 -0.392 57.220 1.00 22.15 AA 30.651 -0.392 57.220 1.00 32.07 AA 28.376 -2.980 57.107 1.00 33.19 AA 27.975 -2.281 56.057 1.00 33.58 AA 27.975 -2.281 56.057 1.00 33.58 AA 27.975 -2.883 59.481 1.00 33.696 AA 27.975 -2.883 59.481 1.00 33.19 AA 27.930 -2.883 59.481 1.00 33.19 AA 27.930 -2.883 59.481 1.00 33.19 AA 27.352 -4.917 58.155 1.00 37.34 AA 28.803 -2.019 59.546 1.00 31.63 AA 29.154 -4.917 58.155 1.00 37.34 AA 29.154 -4.917 58.155 1.00 37.34 AA 29.154 -4.917 58.155 1.00 37.34 AA 29.131 -3.567 62.266 1.00 26.64 AA 28.977 -4.954 62.893 1.00 38.16 AA 29.131 -3.567 62.266 1.00 27.48 AA 29.131 -3.567 62.266 1.00 27.48 AA 29.360 -6.069 62.091 1.00 32.77 AA 28.807 -8.825 54.171 1.00 33.11 AA 28.607 -8.825 54.171 1.00 33.15 AA 28.7382 -1.496 61.770 1.00 33.13 AA 28.607 -8.825 54.171 1.00 33.15 AA 28.318 -5.160 64.047 1.00 33.11 AA 28.607 -8.825 64.171 1.00 34.58 AA 29.336 -6.440 64.512 1.00 34.25 AA 29.336 -6.440 64.512 1.00 34.25 AA 29.337 -2.846 62.261 1.00 34.51 AA 28.337 -2.866 62.121 1.00 37.48 AA 28.338 -5.160 64.047 1.00 39.97 AA 28.339 -6.286 62.277 1.00 34.25 AA 28.318 -5.160 64.047 1.00 31.18 AA 28.607 -8.656 62.727 1.00 34.25 AA 28.339 -0.286 62.121 1.00 31.85 AA 28.349 -7.543 62.266 1.00 26.662 AA 28.357 -0.286 62.277 1.00 34.25 AA 28.360 -6.669 62.091 1.00 34.25 AA 28.300 -6.660 62.570 0.00 34.25 AA 28.318 -5.160 64.047 1.00 33.175 AA 28.32 -1.496 61.770 1.00 31.93 AA 28.5584 -0.117 66.588 1.00 23.75 AA 28.6669 -5.5
ATOM 1050 CD ARG 108 ATOM 1051 NE ARG 108 ATOM 1053 CZ ARG 108 ATOM 1053 CZ ARG 108 ATOM 1057 NH2 ARG 108 ATOM 1060 C ARG 108 ATOM 1061 O ARG 108 ATOM 1061 O ARG 108 ATOM 1062 N GLY 109 ATOM 1066 C GLY 109 M 1066 C GLY 109 M 1066 C GLY 109 M 1067 N ALA 110 ATOM 1070 CB ALA 110 ATOM 1071 C ALA 110 ATOM 1071 C ALA 110 ATOM 1075 CA ILE 111 ATOM 1075 CA ILE 111 ATOM 1076 CB ILE 111 ATOM 1077 CGZ ILE 111 ATOM 1078 CGI ILE 111 ATOM 1078 CGI ILE 111 ATOM 1080 C ILE 111 ATOM 1080 C ILE 111 ATOM 1081 O ILE 111 ATOM 1082 N ARG 112 ATOM 1084 CA ARG 112 ATOM 1085 CB ARG 112 ATOM 1086 CG ARG 112 ATOM 1087 CD ARG 112 ATOM 1087 CD ARG 112 ATOM 1090 CZ ARG 112 ATOM 1090 CZ ARG 112 ATOM 1091 NH1 ARG 112 ATOM 1097 C ARG 112 ATOM 1098 O ARG 112 ATOM 1097 C ARG 112 ATOM 1098 O ARG 112 ATOM 1099 C ILE 113 ATOM 1090 CZ ARG 114 ATOM 1091 NH1 ARG 112 ATOM 1091 NH1 ARG 112 ATOM 1092 O ARG 112 ATOM 1093 O ARG 112 ATOM 1094 O ARG 112 ATOM 1096 O ARG 112 ATOM 1097 C ARG 112 ATOM 1098 O ARG 112 ATOM 1099 O ILE 113 ATOM 1104 CCI ILE 113 ATOM 1105 CDI ILE 113 ATOM 1106 C ILE 113 ATOM 1107 O ILE 113 ATOM 1108 N GLU 114 ATOM 1116 C GLU 114 ATOM 1116 C GLU 114 ATOM 1117 O GLU 114 ATOM 1116 C GLU 114 ATOM 1116 C GLU 114 ATOM 1117 O GLU 114 ATOM 1118 N LYS 115 ATOM 1122 CG LYS 115 ATOM 1123 CD LYS 115 ATOM 1124 CE LYS 115 ATOM 1125 NZ LYS 115	33.935 22.322 65.272 1.00 19.02  33.532 23.268 64.238 1.00 25.61  34.268 23.562 63.1.59 1.00 29.99  35.445 22.977 62.999 1.00 32.55  32.584 18.382 64.791 1.00 23.94  33.427 17.830 64.199 1.00 28.69  31.388 16.724 64.329 1.00 24.68  30.991 18.361 62.982 1.00 24.93  30.285 17.007 63.022 1.00 28.30  30.297 16.356 64.063 1.00 31.84  29.683 16.577 61.906 1.00 28.32  28.934 15.322 61.835 1.00 23.60  27.573 15.585 61.269 1.00 19.14  29.628 14.307 60.974 1.00 22.79  30.707 14.543 60.467 1.00 26.36  29.554 12.123 59.988 1.00 19.51  29.554 12.123 59.988 1.00 19.28  29.512 10.816 60.758 1.00 16.95  29.821 11.080 62.219 1.00 16.95  29.821 10.200 60.659 1.00 16.95  28.138 8.756 61.059 1.00 18.27  27.699 12.497 58.596 1.00 21.80  27.699 12.497 58.596 1.00 21.80  27.699 12.497 58.596 1.00 22.65  28.976 11.643 54.004 1.00 22.65  28.976 11.643 54.004 1.00 21.34  28.781 12.820 53.155 1.00 23.45  27.949 12.552 52.011 1.00 27.02  28.976 11.643 54.004 1.00 21.34  28.781 12.820 53.155 1.00 24.45  27.695 13.460 51.075 1.00 33.45  28.217 14.676 51.77 1.00 35.06  28.221 14.676 51.77 1.00 35.86  29.061 9.730 55.897 1.00 24.85  30.922 13.160 50.339 1.00 27.02  28.964 6.735 56.540 1.00 27.02  28.976 1.643 54.004 1.00 27.22  28.467 7.370 55.677 1.00 27.22  28.467 7.370 55.677 1.00 27.22  28.467 7.370 55.677 1.00 27.22  28.467 7.370 55.677 1.00 27.22  28.467 7.370 55.677 1.00 27.22  28.467 7.370 55.677 1.00 27.49  27.585 7.067 54.482 1.00 29.19  26.424 6.723 54.653 1.00 31.01  27.585 7.067 54.482 1.00 29.19  26.584 10.563 50.492 1.00 22.36  27.585 7.067 54.482 1.00 29.19  26.584 10.563 50.492 1.00 32.27  27.128 8.389 50.001 0.00 27.17  29.159 6.104 50.645 1.00 31.01  27.5979 6.104 50.645 1.00 31.01  27.5979 6.104 50.645 1.00 31.01  27.991 5.591 6.665 40.19 1.00 27.22  28.166 6.905 50.398 1.00 27.17  29.159 6.104 50.645 1.00 31.01  27.184 55.54 50.993 1.00 27.17  29.159 6.104 50.645 1.00 31.01  27.991 5.591 6.666 699 6.00 27.11  29.159 6.104 50.645 1.00 31.01  27.184 5.54 6.998 46.972 1.00 23.36	AAAA ATOM 1217 CS1 VAL 125 AAAA ATOM 1218 CS2 VAL 125 AAAA ATOM 1219 C VAL 125 AAAA ATOM 1220 O VAL 125 AAAA ATOM 1221 N ASP 126 AAAA ATOM 1221 N ASP 126 AAAA ATOM 1223 CA ASP 126 AAAA ATOM 1223 CA ASP 126 AAAA ATOM 1225 CG ASP 126 AAAA ATOM 1226 CD1 ASP 126 AAAA ATOM 1226 CD1 ASP 126 AAAA ATOM 1228 C ASP 126 AAAA ATOM 1228 C ASP 126 AAAA ATOM 1229 O ASP 126 AAAA ATOM 1229 O ASP 126 AAAA ATOM 1230 N TRP 127 AAAA ATOM 1231 CB TRP 127 AAAA ATOM 1231 CB TRP 127 AAAA ATOM 1233 CB TRP 127 AAAA ATOM 1235 CD2 TRP 127 AAAA ATOM 1236 CE2 TRP 127 AAAA ATOM 1237 CE3 TRP 127 AAAA ATOM 1238 CD1 TRP 127 AAAA ATOM 1238 CD1 TRP 127 AAAA ATOM 1238 CD1 TRP 127 AAAA ATOM 1241 CZ2 TRP 127 AAAA ATOM 1241 CZ2 TRP 127 AAAA ATOM 1241 CZ2 TRP 127 AAAA ATOM 1242 CZ3 TRP 127 AAAA ATOM 1242 CZ3 TRP 127 AAAA ATOM 1240 CG TRP 127 AAAA ATOM 1241 CZ2 TRP 127 AAAA ATOM 1242 CZ3 TRP 127 AAAA ATOM 1240 CG TRP 127 AAAA ATOM 1241 CZ2 TRP 127 AAAA ATOM 1242 CZ3 TRP 127 AAAA ATOM 1241 CZ2 TRP 127 AAAA ATOM 1242 CZ3 TRP 127 AAAA ATOM 1243 CH2 TRP 127 AAAA ATOM 1244 C TRP 127 AAAA ATOM 1245 O TRP 127 AAAA ATOM 1246 C TRP 127 AAAA ATOM 1245 O TRP 127 AAAA ATOM 1246 C TRP 127 AAAA ATOM 1248 CA SER 128 AAAA ATOM 1249 CB SER 128 AAAA ATOM 1260 CG SER	28. 602 3.609 66.620 1.00 20.52 AAA 28.725 1.496 65.462 1.00 20.22 AAA 25.029 3.695 66.669 1.00 29.23 AAA 25.029 3.695 66.669 1.00 29.23 AAA 25.029 3.695 66.669 1.00 29.23 AAA 25.029 3.695 68.588 1.00 28.67 AAA 25.029 3.695 69.106 1.00 27.48 AAA 25.029 3.695 69.106 1.00 27.48 AAA 25.029 3.695 69.106 1.00 29.26 AAA 24.905 4.435 70.572 1.00 29.26 AAA 23.757 5.320 71.073 1.00 29.89 AAA 26.095 6.025 68.978 1.00 30.01 AAA 26.095 6.025 68.978 1.00 30.01 AAA 26.111 6.596 67.776 1.00 27.25 AAA 26.695 6.025 68.978 1.00 30.01 AAA 26.111 6.596 67.776 1.00 23.75 AAA 27.124 7.316 64.969 1.00 25.40 AAA 28.477 7.067 64.529 1.00 23.75 AAA 28.477 7.067 64.529 1.00 23.72 AAA 28.412 6.084 63.514 1.00 25.60 AAA 29.729 7.584 64.900 1.00 25.60 AAA 27.040 10.03 64.255 1.00 23.74 AAA 27.095 5.752 63.344 1.00 25.60 AAA 29.551 5.603 62.862 1.00 24.56 AAA 29.551 5.603 62.862 1.00 24.56 AAA 27.040 10.037 68.424 1.00 25.08 AAA 27.040 10.037 68.424 1.00 25.50 AAA 27.040 10.037 68.424 1.00 25.08 AAA 27.040 10.037 68.424 1.00 25.09 AAA 27.040 10.037 68.424 1.00 25.09 AAA 28.803 25.646 69.679 1.00 25.03 AAA 29.189 9.806 71.289 1.00 25.03 AAA 29.189 9.806 71.289 1.00 25.03 AAA 20.288 809 2.701 71.402 1.00 22.49 AAA 20.294 81 10.116 69.768 1.00 26.10 AAA 20.294 81 10.116 69.768 1.00 26.10 AAA 20.294 81 1

ATOM 1285 CG ASP 132 ATOM 1286 OD1 ASP 132 1287 OD2 ASP 132 1288 C ASP 132 A. A 1289 O ASP 132	27.203 17.299 71.683 1.00 40.20 26.867 16.379 72.461 1.00 46.09 27.531 18.454 72.055 1.00 44.38 25.570 15.814 68.876 1.00 29.06 24.460 15.362 69.093 1.00 34.86	AAAA ATGM 1445 C ASP 150 AAAA ATGM 1446 O ASP 150 AAAA ATGM 1447 N LEU 151 AAAA ATGM 1449 CA LEU 151 AAAA ATGM 1450 CB LEU 151	24.550 -8.922 63.833 1.00 41.30 AAAA 23.458 -8.771 64.376 1.00 44.58 AAAA 25.536 -9.597 64.398 1.00 37.30 AAAA 25.359 -10.181 65.699 1.00 35.09 AAAA 25.407 -11.702 65.571 1.00 38.32 AAAA
ATOM 1290 N ALA 133 ATOM 1292 CA ALA 133 ATOM 1293 CB ALA 133 ATOM 1294 C ALA 133 ATOM 1295 O ALA 133 ATOM 1296 N VAL 134 ATOM 1298 CA VAL 134 ATOM 1299 CB VAL 134	25.913 16.511 67.792 1.00 26.57 25.060 16.891 66.665 1.00 22.34 25.767 17.973 65.893 1.00 22.29 24.683 15.764 65.703 1.00 22.13 24.814 15.896 64.478 1.00 20.05 24.194 14.667 66.252 1.00 22.27 23.813 13.524 65.448 1.00 24.74 23.360 12.371 66.353 1.00 23.00	AAA ATOM 1451 OG LEU 151 AAAA ATOM 1453 C LEU 151 AAAA ATOM 1454 O LEU 151 AAAA ATOM 1455 N CYS 152 AAAA ATOM 1457 CA CYS 152 AAAA ATOM 1458 C CYS 152 AAAA ATOM 1459 O CYS 152 AAAA ATOM 1459 O CYS 152 AAAA ATOM 1460 CB CYS 152	24.168 -12.207 65.097 1.00 36.38 AAA 26.452 -9.655 66.624 1.00 34.61 AAA 27.640 -9.885 66.409 1.00 33.41 AAA 26.045 -8.937 67.659 1.00 35.32 AAA 27.006 -8.365 68.593 1.00 40.29 AAA 27.570 -9.463 69.480 1.00 43.08 AAA 26.977 -10.542 69.568 1.00 45.04 AAA 26.342 -7.288 69.473 1.00 37.72 AAAA
ATOM 1300 CG1 VAL 134 ATOM 1301 CG2 VAL 134 ATOM 1302 C VAL 134 ATOM 1303 O VAL 134	22.194 12.804 67.154 1.00 27.35 22.996 11.152 65.547 1.00 29.73 22.699 13.884 64.478 1.00 29.00 22.585 13.283 62.415 1.00 32.69 21.880 14.869 64.824 1.00 29.17	AAA ATOM 1461 SG CYS 152  AAA ATOM 1462 N PRO 153  AAA ATOM 1463 CD PRO 153  AAA ATOM 1464 CA PRO 153  AAAA ATOM 1465 CB PRO 153	26.098 -5.631 68.736 1.00 17.05 AAAA 28.739 -9.216 70.121 1.00 45.27 AAAA 29.565 -8.005 69.975 1.00 41.44 AAAA 29.381 -10.190 71.012 1.00 47.26 AAAA
ATOM 1304 N SER 135 ATOM 1306 CA SER 135 ATOM 1307 CB SER 135 ATOM 1310 C SER 135 ATOM 1311 O SER 135 ATOM 1311 O SER 135 ATOM 1311 O SER 135 ATOM 1312 N ASN 136 ATOM 1314 CA ASN 136 ATOM 1315 CB ASN 136 ATOM 1316 CG ASN 136 ATOM 1317 OD1 ASN 136 ATOM 1318 ND2 ASN 136 ATOM 1321 C ASN 136 ATOM 1322 O ASN 136 ATOM 1322 O ASN 136 ATOM 1322 C ASN 137 ATOM 1325 CA ASN 137 ATOM 1326 CB ASN 137 ATOM 1326 CB ASN 137 ATOM 1327 CG ASN 137 ATOM 1328 OD1 ASN 137 ATOM 1329 ND2 ASN 137 ATOM 1332 C ASN 137 ATOM 1333 O ASN 137 ATOM 1334 N TYR 138 ATOM 1336 CA TYR 138 ATOM 1337 CB TYR 138 ATOM 1338 CG TYR 138 ATOM 1339 CD1 TYR 138 ATOM 1341 CD2 TYR 138 ATOM 1342 CE2 TYR 138 ATOM 1342 CE2 TYR 138 ATOM 1344 OH TYR 138 ATOM 1345 C TYR 138 ATOM 1346 C TYR 138 ATOM 1347 O TYR 138 ATOM 1348 N ILE 139 ATOM 1348 N ILE 139 ATOM 1350 CA ILE 139 ATOM 1351 CB ILE 139 ATOM 1352 CG2 ILE 139 ATOM 1355 C ILE 139 ATOM 1355 C ILE 139 ATOM 1356 O ILE 139 ATOM 1356 O ILE 139 ATOM 1356 O ILE 139 ATOM 1357 N VAL 140	20.796       15.257       63.950       1.00       29.17         20.111       16.477       64.504       1.00       27.10         21.062       17.327       65.070       1.00       23.39         21.232       15.552       62.511       1.00       34.63         20.404       15.509       61.586       1.00       36.08         22.506       15.866       62.272       1.00       38.35         22.847       16.132       60.82       1.00       39.46         23.071       17.638       60.650       1.00       43.49         24.275       18.155       61.314       1.00       49.28         24.266       18.420       62.514       1.00       51.34         25.349       18.334       60.533       1.00       58.07         23.897       15.261       60.189       1.00       35.57         24.619       15.688       59.273       1.00       35.22         23.894       14.002       60.613       1.00       35.22         23.894       14.002       60.613       1.00       27.82         24.732       11.747       60.947       1.00       20.95	AAAA ATOM 1465 CB PRO 153 AAAA ATOM 1466 CG PRO 153 AAAA ATOM 1467 C PRO 153 AAAA ATOM 1468 O PRO 153 AAAA ATOM 1469 N GLY 154 AAAA ATOM 1471 CA GLY 154 AAAA ATOM 1472 C GLY 154 AAAA ATOM 1473 O GLY 154 AAAA ATOM 1474 N THR 155 AAAA ATOM 1476 CA THR 155 AAAA ATOM 1477 CB THR 155 AAAA ATOM 1478 OG1 THR 155 AAAA ATOM 1480 CG2 THR 155 AAAA ATOM 1481 C THR 155 AAAA ATOM 1482 O THR 155 AAAA ATOM 1483 N MET 156 AAAA ATOM 1485 CA MET 156 AAAA ATOM 1485 CA MET 156 AAAA ATOM 1486 CB MET 156 AAAA ATOM 1488 SD MET 156 AAAA ATOM 1488 SD MET 156 AAAA ATOM 1488 SD MET 156 AAAA ATOM 1489 CE MET 156 AAAA ATOM 1490 C MET 156 AAAA ATOM 1490 C MET 156 AAAA ATOM 1491 O MET 156 AAAA ATOM 1492 N GLU 157 AAAA ATOM 1494 CA GLU 157 AAAA ATOM 1496 C GLU 157 AAAA ATOM 1497 O GLU 157 AAAA ATOM 1498 N GLU 157 AAAA ATOM 1498 N GLU 157 AAAA ATOM 1498 N GLU 158 AAAA ATOM 1501 CB GLU 158 AAAA ATOM 1501 CB GLU 158 AAAA ATOM 1500 CA GLU 158	30.591 -9.439 71.562 1.00 46.10 AAAA 10.894 -8.438 70.530 1.00 44.99 28.442 -10.694 72.113 1.00 50.81 AAAA 27.825 -9.923 72.865 1.00 47.27 AAAA 28.339 -12.015 72.167 1.00 57.66 AAAA 27.477 -12.680 73.125 1.00 67.39 AAAA 25.998 -14.517 72.652 1.00 75.86 AAAA 25.998 -14.517 72.652 1.00 75.86 AAAA 25.5992 -12.553 71.612 1.00 75.09 AAAA 24.400 -12.955 70.874 1.00 76.56 AAAA 24.209 -12.007 69.709 1.00 74.93 AAAA 24.209 -12.007 69.709 1.00 74.93 AAAA 22.932 -12.329 68.965 1.00 74.24 AAAA 22.932 -12.329 68.965 1.00 74.24 AAAA 23.234 -15.039 70.602 1.00 80.09 AAAA 23.234 -15.039 70.602 1.00 80.09 AAAA 25.191 -16.219 69.038 1.00 89.85 AAAA 25.191 -16.219 69.038 1.00 89.85 AAAA 26.072 -16.305 67.777 1.00 93.38 AAAA 26.072 -16.305 67.777 1.00 93.38 AAAA 26.072 -16.732 65.287 1.00100.00 AAAA 22.937 -16.735 65.851 1.00 90.63 AAAA 24.697 -16.732 65.287 1.00100.00 AAAA 25.611 -17.282 70.053 1.00 90.29 AAAA 25.611 -17.282 70.053 1.00 90.31 AAAA 25.611 -17.282 70.053 1.00 90.31 AAAA 25.918 -16.820 71.258 1.00 90.33 AAAA 26.035 -17.690 72.360 1.00 87.84 AAAA 25.320 -17.500 73.512 1.00 87.84 AAAA 25.320 -17.500 73.512 1.00 87.84 AAAA 26.032 -17.500 73.512 1.00 87.84 AAAA 26.032 -17.500 73.512 1.00 88.22 AAAA 26.032 -17.500 73.512 1.00 88.22 AAAA 26.032 -17.500 73.512 1.00 88.22 AAAA 26.032 -17.500 73.512 1.00 87.84 AAAA 27.735 -17.337 72.819 1.00 91.16 AAAA 27.735 -17.337 72.819 1.00 99.50 AAAA 28.3047 -16.845 74.135 1.00 88.22 AAAA 29.684 -17.845 74.202 1.00 94.30 AAAA 20.684 -17.845 74.202 1.00 94.30 AAAA 21.660 -18.479 75.663 1.00 77.31 AAAA 22.5061 -13.661 77.990 72.865 1.00 97.31 AAAA 23.616 -16.384 75.460 1.00 79.85 AAAA 23.616 -16.384 75.460 1.00 77.31 AAAA 24.660 -15.574 75.356 1.00 77.31 AAAA 25.861 -13.665 75.997 1.00 76.20 AAAA 24.660 -15.675 76.893 1.00 77.31 AAAA 25.861 -13.665 75.997 1.00 74.53 AAAA 25.861 -13.665 75.997 1.00 74.53 AAAA 26.572 -15.661 77.995 76.497 7.00 76.20 AAAA
ATOM 1357 N VAL 140 ATOM 1359 CA VAL 140 ATOM 1360 CB VAL 140 ATOM 1361 CG1 VAL 140 ATOM 1362 CG2 VAL 140 ATOM 1363 C VAL 140 ATOM 1364 O VAL 140	22,578 6.560 52.694 1.00 26.60 21,881 7.660 51.875 1.00 27.22 22,850 8,255 50.882 1.00 23.87 20,616 7.087 51.169 1.00 26.11 23,392 5.694 51.754 1.00 30.61 24,579 5.945 51.567 1.00 34.21	AAA ATOM 1515 CD PRO 160 AAA ATOM 1516 CA PRO 160 AAA ATOM 1517 CB PRO 160 AAA ATOM 1518 CG PRO 160 AAA ATOM 1519 C PRO 160 AAA ATOM 1520 O PRO 160	23.560 -12.800 76.372 1.00 72.38 AAAA 25.213 -11.269 75.476 1.00 69.47 AAAA 23.843 -10.597 75.562 1.00 69.23 AAAA 23.108 -11.381 76.566 1.00 70.96 AAAA 26.272 -10.481 76.222 1.00 66.84 AAAA 26.306 -10.439 77.458 1.00 66.35 AAAA
ATOM 1365 N GLY 141 ATOM 1365 C GLY 141 ATOM 1369 O GLY 141 ATOM 1370 N ASN 142 ATOM 1371 CB ASN 142 ATOM 1373 CB ASN 142 ATOM 1375 CD1 ASN 142 ATOM 1376 ND2 ASN 142 ATOM 1380 O ASN 142 ATOM 1381 N LYS 143 ATOM 1381 N LYS 143 ATOM 1383 CA LYS 143 ATOM 1384 CB LYS 143 ATOM 1388 NZ LYS 143 ATOM 1388 NZ LYS 143 ATOM 1398 CC LYS 143 ATOM 1399 C LYS 143 ATOM 1399 C D PRO 144 ATOM 1396 CA PRO 144 ATOM 1397 CB PRO 144 ATOM 1396 CA PRO 144 ATOM 1397 CB PRO 144 ATOM 1399 C PRO 144 ATOM 1399 C PRO 144 ATOM 1390 CG PRO 144 ATOM 1400 O PRO 144 ATOM 1401 N PRO 145 ATOM 1403 CA PRO 145 ATOM 1403 CA PRO 145 ATOM 1404 CB PRO 145 ATOM 1405 CG PRO 145 ATOM 1406 C PRO 145 ATOM 1407 O PRO 145 ATOM 1408 N LYS 146 ATOM 1409 C G GLY 147 ATOM 1410 CA LYS 146 ATOM 1411 CB LYS 146 ATOM 1412 CG LYS 146 ATOM 1412 CG LYS 146 ATOM 1415 O LYS 146 ATOM 1416 N GLU 147 ATOM 1418 CA GLU 147 ATOM 1418 CA GLU 147 ATOM 1419 CB GLU 147 ATOM 1418 CA GLU 147 ATOM 1420 CC GLU 147 ATOM 1418 CA GLU 147 ATOM 1418 CA GLU 147 ATOM 1418 CA GLU 147 ATOM 1420 CC GLU 147 ATOM 1421 CB GLU 147 ATOM 1422 CG GLU 147 ATOM 1423 CC GLU 147 ATOM 1424 C GLU 147 ATOM 1425 CG GLU 147 ATOM 1426 C GLU 147 ATOM 1427 CC GLU 147 ATOM 1428 CA CYS 148 ATOM 1439 CG PS 148 ATOM 1441 CB ASP 150 ATOM 1431 CB CYS 148 ATOM 1442 CG GLU 147 ATOM 1423 CC GLU 147 ATOM 1424 C GLU 147 ATOM 1425 CG GLU 147 ATOM 1426 C GLU 147 ATOM 1427 CG GLU 147 ATOM 1428 CA CYS 148 ATOM 1430 C GYS 148 ATOM 1441 CB ASP 150	22.751	AAA ATOM 1521 N MET 161 AAA ATOM 1524 CB MET 161 AAA ATOM 1524 CB MET 161 AAA ATOM 1526 CG MET 161 AAA ATOM 1526 CG MET 161 AAA ATOM 1526 CG MET 161 AAA ATOM 1527 CE MET 161 AAA ATOM 1528 C MET 161 AAA ATOM 1528 C MET 161 AAA ATOM 1529 O MET 161 AAA ATOM 1529 O MET 161 AAA ATOM 1530 N CYS 162 AAA ATOM 1531 C CYS 162 AAA ATOM 1531 C CYS 162 AAA ATOM 1533 C CYS 162 AAA ATOM 1533 C CYS 162 AAA ATOM 1533 C CYS 162 AAA ATOM 1536 CG CYS 162 AAA ATOM 1536 CG CYS 162 AAA ATOM 1537 N GLU 163 AAA ATOM 1538 CA GLU 163 AAA ATOM 1540 CB GLU 163 AAA ATOM 1540 CB GLU 163 AAA ATOM 1541 CG GLU 163 AAA ATOM 1542 CD GLU 163 AAA ATOM 1544 OE2 GLU 163 AAA ATOM 1545 C GLU 163 AAA ATOM 1547 N LYS 164 AAA ATOM 1546 O GLU 163 AAA ATOM 1547 N LYS 164 AAA ATOM 1547 N LYS 164 AAA ATOM 1549 CA LYS 164 AAA ATOM 1550 CB LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1550 CB LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1550 CB LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1550 CB LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1550 CB LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1550 CB LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1551 CG LYS 164 AAA ATOM 1556 CB LYS 164 AAA ATOM 1557 CB LYS 164 AAA ATOM 1556 CB LYS 164 AAA ATOM 1560 N THR 165 AAA ATOM 1560 N THR 165 AAA ATOM 1560 CB LYS 164 AAA ATOM 1560 CB LYS 164 AAA ATOM 1560 N THR 165 AAA ATOM 1560 N THR 165 AAA ATOM 1560 N THR 165 AAA ATOM 1560 N THR 166 AAA ATOM 1560 N	27.111 -9.814 75.443 1.00 63.26 AAAA 28.180 -9.028 76.005 1.00 60.06 AAAA 29.448 -9.266 75.192 1.00 61.41 AAAA 30.084 -10.599 75.458 1.00 63.14 AAAA 30.315 -10.833 77.229 1.00 67.62 AAAA 31.591 -12.057 77.212 1.00 58.22 AAAA 27.919 -7.523 76.114 1.00 58.22 AAAA 28.013 -6.960 77.213 1.00 53.07 AAAA 27.575 -6.883 74.990 1.00 53.00 AAAA 27.331 -5.427 74.938 1.00 47.97 AAAA 26.175 -4.829 75.590 1.00 47.83 AAAA 25.187 -5.513 75.801 1.00 51.51 AAAA 27.404 -4.981 73.503 1.00 45.58 AAAA 28.346 -6.145 72.496 1.00 47.42 AAAA 26.249 -1.536 75.897 1.00 45.59 AAAA 26.778 -1.755 78.156 1.00 57.92 AAAA 26.5778 -1.755 78.156 1.00 57.92 AAAA 26.778 -1.755 78.156 1.00 57.92 AAAA 26.778 -1.755 78.156 1.00 57.92 AAAA 26.679 -1.080 79.509 1.00 64.85 AAAA 26.461 -1.824 80.510 1.00 69.44 AAAA 24.061 -2.456 75.503 1.00 44.29 AAAA 24.061 -2.456 75.503 1.00 44.29 AAAA 24.061 -2.456 75.503 1.00 44.29 AAAA 24.061 -2.456 75.503 1.00 44.99 AAAA 24.061 -2.787 77.982 1.00 49.91 AAAA 24.061 -2.787 77.982 1.00 49.91 AAAA 24.061 -2.787 77.982 1.00 49.91 AAAA 24.061 -2.787 77.78 77.982 1.00 49.91 AAAA 24.061 -2.550 77.77 1.00 1.00 1.00 47.89 AAAA 24.061 -2.456 75.503 1.00 48.92 AAAA 24.061 -2.550 77.77 1.00 1.00 38.39 AAAA 24.061 -2.550 77.77 1.00 38.39 AAAA 24.061 -2.456 77.509 1.00 48.91 AAAA 24.061 -2.456 77.509 1.00 48.91 AAAA 24.061 -2.456 77.509 1.00 49.91 AAAA 24.061 -2.456 77.509 1.00 38.39 AAAA 24.787 97.982 1.00 38.99 AAAA 24.787 97.982 1.00 38.91 AAAA 24.787 97.982 1.00 38.99 AAAA 24.787 97.982 1.00 38.99 AAAA 24.787 97.982 97.982 1.00 38.99 AAAA 24.787 97.986 97.982 97.982 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.992 97.99

ATOM 1603 OD1 ASN 169 AT 1604 ND2 ASN 169 1607 C ASN 169 1608 O ASN 169 A. 1609 N GLU 170 ATOM 1611 CA GLU 170 ATOM 1611 CA GLU 170 ATOM 1613 CG GLU 170 ATOM 1614 CD GLU 170 ATOM 1615 OEI GLU 170 ATOM 1616 OE2 GLU 170 ATOM 1617 C GLU 170 ATOM 1618 O GLU 170 ATOM 1619 N TYR 171 ATOM 1621 CA TYR 171 ATOM 1622 CB TYR 171 ATOM 1622 CB TYR 171 ATOM 1624 CD1 TYR 171 ATOM 1625 CE1 TYR 171 ATOM 1626 CD2 TYR 171 ATOM 1627 CE2 TYR 171 ATOM 1628 CZ TYR 171 ATOM 1631 C TYR 171 ATOM 1631 C TYR 171 ATOM 1632 O TYR 171 ATOM 1633 O TYR 171 ATOM 1635 CA ASN 172 ATOM 1635 CA ASN 172 ATOM 1635 CA ASN 172 ATOM 1636 CB ASN 172 ATOM 1638 OD1 ASN 172 ATOM 1638 OD1 ASN 172 ATOM 1640 CB ASN 172 ATOM 1640 CB TYR 173 ATOM 1650 CB TYR 173 ATOM 1650 CB TYR 173 ATOM 1640 CB TYR 173 ATOM 1650 CB TYR 173 ATOM 1650 CB TYR 173 ATOM 1660 CB ASN 172 ATOM 1661 CB ASN 172 ATOM 1663 CB TYR 173 ATOM 1664 CA TYR 173 ATOM 1664 CA TYR 173 ATOM 1665 CB TYR 173 ATOM 1664 CB TYR 173 ATOM 1665 CB TYR 173 ATOM 1665 CB TYR 173 ATOM 1666 CA ARG 174 ATOM 1665 CB TYR 173 ATOM 1666 CB ARG 174 ATOM 1666 CB ARG 174 ATOM 1667 CB ARG 174 ATOM 1668 CB CYS 175 ATOM 1668 CB TYR 173 ATOM 1666 CB TYR 173 ATOM 1666 CB CYS 175 ATOM 1668 CB TRP 176	10.912	AAAA ATOM 1775 O LYS 183 AAAA ATOM 1776 N MET 184 AAAA ATOM 1778 CA MET 184 AAAA ATOM 1778 CB MET 184 AAAA ATOM 1780 CG MET 184 AAAA ATOM 1781 SD MET 184 AAAA ATOM 1781 SD MET 184 AAAA ATOM 1782 CE MET 184 AAAA ATOM 1782 CE MET 184 AAAA ATOM 1783 C MET 184 AAAA ATOM 1783 C MET 184 AAAA ATOM 1783 C MET 184 AAAA ATOM 1785 N CYS 185 AAAA ATOM 1787 CA CYS 185 AAAA ATOM 1789 C CYS 185 AAAA ATOM 1789 C CYS 185 AAAA ATOM 1790 CB CYS 185 AAAA ATOM 1790 CB CYS 185 AAAA ATOM 1791 SG CYS 185 AAAA ATOM 1792 N DRO 186 AAAA ATOM 1792 N DRO 186 AAAA ATOM 1795 CB PRO 186 AAAA ATOM 1796 CG PRO 186 AAAA ATOM 1797 C PRO 186 AAAA ATOM 1798 O PRO 186 AAAA ATOM 1799 N SER 187 AAAA ATOM 1801 CA SER 187 AAAA ATOM 1801 CA SER 187 AAAA ATOM 1802 CB SER 187 AAAA ATOM 1805 C SER 187 AAAA ATOM 1805 C SER 187 AAAA ATOM 1806 O SER 187 AAAA ATOM 1807 N THR 188 AAAA ATOM 1808 C THR 188 AAAA ATOM 1810 CB THR 188 AAAA ATOM 1811 CG1 THR 188 AAAA ATOM 1812 CG YS 189 AAAA ATOM 1813 CG2 THR 188 AAAA ATOM 1814 C THR 188 AAAA ATOM 1815 C THR 188 AAAA ATOM 1810 CB THR 189 AAAA ATOM 1	32.015
ATOM 1688 CE2 TRP 176 ATOM 1688 CE2 TRP 176 ATOM 1688 CE2 TRP 176 ATOM 1689 CE3 TRP 176 ATOM 1690 CD1 TRP 176 ATOM 1691 NE1 TRP 176 ATOM 1691 NE1 TRP 176 ATOM 1693 CZ2 TRP 176 ATOM 1694 CZ3 TRP 176 ATOM 1695 CH2 TRP 176 ATOM 1695 CH2 TRP 176 ATOM 1696 C TRP 176 ATOM 1697 O TRP 176 ATOM 1698 N THR 177 ATOM 1700 CA THR 177 ATOM 1701 CB THR 177 ATOM 1702 CG1 THR 177 ATOM 1705 C THR 177 ATOM 1705 C THR 177 ATOM 1706 O THR 177 ATOM 1707 N THR 178 ATOM 1709 CA THR 178 ATOM 1710 CB THR 178 ATOM 1711 CG1 THR 178 ATOM 1711 CG2 THR 178 ATOM 1713 CG2 THR 178 ATOM 1714 C THR 178 ATOM 1715 O THR 178 ATOM 1716 N ASN 179 ATOM 1716 N ASN 179 ATOM 1717 CG ASN 179 ATOM 1718 CA ASN 179 ATOM 1719 CB ASN 179 ATOM 1720 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1721 CD1 ASN 179 ATOM 1720 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1721 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1721 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1721 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1720 CG ASN 179 ATOM 1721 CG ASN 179 ATOM 1722 ND2 ASN 179 ATOM 1724 C ASG 180 ATOM 1735 CZ ARG 180 ATOM 1736 NH1 ARG 180 ATOM 1731 CG ARG 180 ATOM 1733 NE ARG 180 ATOM 1734 C ARG 180 ATOM 1735 CZ ARG 180 ATOM 1736 NH1 ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1737 C ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 CD ARG 180 ATOM 1737 N CR BRG 180 ATOM 1736 CD ARG 180 ATOM 1737 N CR BRG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 CD ARG 180 ATOM 1736 CD ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 CD ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 CD ARG 180 ATOM 1736 CD ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 CD ARG 180 ATOM 1736 NH2 ARG 180 ATOM 1736 CD ARG 180 ATOM 1736 NH2 ARG	32.534	AAAA ATOM 1860 CA ALA 193 AAAA ATOM 1861 CB ALA 193 AAAA ATOM 1862 C ALA 193 AAAA ATOM 1863 O ALA 193 AAAA ATOM 1866 N CYS 194 AAAA ATOM 1866 CA CYS 194 AAAA ATOM 1866 CA CYS 194 AAAA ATOM 1866 CA CYS 194 AAAA ATOM 1867 C CYS 194 AAAA ATOM 1868 O CYS 194 AAAA ATOM 1869 CB CYS 194 AAAA ATOM 1870 SG CYS 194 AAAA ATOM 1871 N THR 195 AAAA ATOM 1873 CA THR 195 AAAA ATOM 1873 CA THR 195 AAAA ATOM 1877 CG2 THR 195 AAAA ATOM 1877 CG2 THR 195 AAAA ATOM 1878 C THR 195 AAAA ATOM 1878 C THR 195 AAAA ATOM 1878 C THR 195 AAAA ATOM 1882 CA GLU 196 AAAA ATOM 1882 CA GLU 196 AAAA ATOM 1885 CD GLU 196 AAAA ATOM 1885 CD GLU 196 AAAA ATOM 1885 CD GLU 196 AAAA ATOM 1886 OE1 GLU 196 AAAA ATOM 1887 OE2 GLU 196 AAAA ATOM 1889 O GLU 196 AAAA ATOM 1889 O GLU 196 AAAA ATOM 1889 O GLU 196 AAAA ATOM 1899 C ASN 197 AAAA ATOM 1891 CB ASN 197 AAAA ATOM 1890 N ASN 197 AAAA ATOM 1891 CB ASN 197 AAAA ATOM 1890 C ASN 197 AAAA ATOM 1900 C ASN 197 AAAA ATOM 1900 C ASN 198 AAAA ATOM 1900 C ASN 1	16.022 9.481 80.293 1.00 34.93 AAAA 17.220 9.828 79.447 1.00 33.97 AAAA 36.416 8.497 81.358 1.00 35.02 AAAA 36.213 8.740 82.531 1.00 32.16 AAAA 36.970 7.168 80.951 1.00 38.60 AAAA 37.366 6.379 81.924 1.00 47.45 AAAA 38.084 4.875 80.208 1.00 49.64 AAAA 36.148 5.816 82.610 1.00 47.45 AAAA 36.148 5.816 82.610 1.00 47.93 AAAA 31.666 6.548 82.610 1.00 47.93 AAAA 31.666 6.61 6.81 81.826 1.00 57.00 AAAA 31.755 3.509 82.037 1.00 53.16 AAAA 31.755 3.509 82.037 1.00 53.16 AAAA 40.401 3.098 83.361 1.00 56.81 AAAA 40.401 3.098 83.361 1.00 56.81 AAAA 41.802 2.549 83.133 1.00 61.17 AAAA 38.980 2.350 81.429 1.00 53.60 AAAA 37.761 2.427 81.288 1.00 55.26 AAAA 39.660 1.289 81.052 1.00 54.62 AAAA 40.814 -0.374 78.647 1.00 59.61 AAAA 40.814 -0.374 78.647 1.00 59.63 AAAA 40.808 0.408 77.450 1.00 64.21 AAAA 40.809 0.408 0.408 1.00 56.03 AAAA 39.877 -0.369 77.460 1.00 62.78 AAAA 40.809 0.408 0.408 1.00 56.03 AAAA 39.877 -0.389 82.819 1.00 55.86 AAAA 38.928 -1.191 77.450 1.00 64.21 AAAA 40.809 0.409 77.460 1.00 62.78 AAAA 40.809 0.409 77.460 1.00 62.78 AAAA 40.809 0.409 77.460 1.00 64.24 AAAA 40.809 0.409 77.460 1.00 65.03 AAAA 40.909 0.400 76.542 1.00 66.04 AAAA 40.909 0.400 76.542 1.00 66.04 AAAA 40.909 0.400 76.542 1.00 66.24 AAAA 40.909 0.400 76.542 1.00 66.27 AAAA 40.909 0.400 76.542 1.00 66.24 AAAA 40.909 0.400 76.542 1.00 66.29 AAAA 40.909 0.400 76.542 1.00 66.29 AAAA 40.909 0.400 76.542 1.00 66.29 AAAA 40.909 0.400 76.542 1.00 66.00 AAAA 40.909 0.900 0

The color of the	ATOM 1940 CG HIS 202 ATOM 1941 CD2 HIS 202 / 1942 ND1 HIS 202 . 1944 CE1 HIS 202 ATOM 1945 NE2 HIS 202 ATOM 1947 C HIS 202 ATOM 1948 O HIS 202 ATOM 1949 N PRO 203 ATOM 1950 CD PRO 203 ATOM 1951 CA PRO 203 ATOM 1951 CA PRO 203 ATOM 1952 CB PRO 203 ATOM 1953 CG PRO 203	42.033 15.187 85.071 1.00 33.88 42.478 15.717 86.230 1.00 34.57 41.801 16.250 84.229 1.00 39.51 42.084 17.379 84.856 1.00 38.52 42.496 17.082 86.073 1.00 34.70 39.332 14.312 84.100 1.00 35.68 39.123 14.067 82.918 1.00 40.37 38.750 15.340 84.710 1.00 34.78 38.883 15.735 86.115 1.00 36.37 37.850 16.233 83.982 1.00 33.97 37.614 17.389 84.954 1.00 32.34 38.582 17.199 86.058 1.00 35.27	AAAA ATOM 2097 CD ARG AAAA ATOM 2098 NE ARG AAAA ATOM 2100 CZ ARG AAAA ATOM 2101 NH1 ARG AAAA ATOM 2104 NH2 ARG AAAA ATOM 2107 C ARG AAAA ATOM 2108 O ARG AAAA ATOM 2109 N HIS AAAA ATOM 2111 CA HIS AAAA ATOM 2112 CB HIS AAAA ATOM 2113 CG HIS AAAA ATOM 2113 CG HIS	222 38.794 19.842 74.961 1.00 41.43 223 40.679 20.036 76.226 1.00 43.22 223 41.062 21.377 75.802 1.00 40.49 223 41.167 22.304 76.996 1.00 40.04 223 39.884 22.489 77.715 1.00 42.81	AAAA AAAA AAAA AAAA AAAA AAAA AAAA
ATOM 1997 CA. CHE 209 14 020 7 7.067 76 809 1.00 41.01 AAAA. ATOM 1156 CE 777 126 127 127 127 127 127 127 127 127 127 127	ATOM 1954 C PRO 203 ATOM 1955 O PRO 203 ATOM 1956 N GLU 204 ATOM 1958 CA GLU 204 ATOM 1959 CB GLU 204 ATOM 1960 CG GLU 204 ATOM 1961 CD GLU 204 ATOM 1961 CD GLU 204 ATOM 1963 OE2 GLU 204 ATOM 1963 OE2 GLU 204 ATOM 1966 C CYS 205 ATOM 1968 CA CYS 205 ATOM 1970 O CYS 205 ATOM 1971 CB CYS 205 ATOM 1971 CB CYS 205 ATOM 1972 SG CYS 205 ATOM 1973 N LEU 206 ATOM 1975 CA LEU 206 ATOM 1976 CB LEU 206 ATOM 1977 CG LEU 206 ATOM 1978 CD1 LEU 206 ATOM 1978 CD1 LEU 206 ATOM 1980 C LEU 206 ATOM 1981 O LEU 206 ATOM 1981 O LEU 206 ATOM 1982 N GLY 207 ATOM 1984 CA GLY 207 ATOM 1985 C GLY 207 ATOM 1985 C GLY 207 ATOM 1986 O GLY 207 ATOM 1987 N SER 208 ATOM 1989 CA SER 208 ATOM 1991 OG SER 208 ATOM 1991 OG SER 208 ATOM 1993 C SER 208 ATOM 1993 C SER 208 ATOM 1994 O SER 208	38.381 16.724 82.642 1.00 34.97 37.614 17.144 81.765 1.00 36.16 39.693 16.676 82.477 1.00 33.97 40.290 17.148 81.242 1.00 31.62 41.545 17.955 81.552 1.00 29.26 41.235 19.253 82.260 1.00 30.14 40.739 20.368 81.331 1.00 34.01 40.657 20.192 80.085 1.00 35.20 40.434 21.445 81.868 1.00 33.07 40.603 16.003 80.299 1.00 30.09 40.585 16.151 79.082 1.00 30.45 40.865 14.835 80.840 1.00 29.97 41.180 13.767 79.931 1.00 32.60 40.046 13.710 78.966 1.00 31.35 38.908 13.801 79.362 1.00 35.49 41.327 12.444 80.662 1.00 33.33 42.981 12.223 81.401 1.00 38.85 40.365 13.637 77.691 1.00 31.01 39.345 13.523 76.677 1.00 30.91 39.346 14.655 75.637 1.00 28.62 38.601 14.521 74.370 1.00 24.69 37.198 14.146 74.723 1.00 23.66 38.580 15.791 73.620 1.00 21.51 39.579 12.154 76.040 1.00 32.93 40.706 11.677 75.931 1.00 35.10 38.510 11.488 75.650 1.00 35.10 38.510 11.488 75.650 1.00 36.31 38.688 10.195 75.033 1.00 38.16 38.752 9.019 75.979 1.00 39.99 37.793 8.239 76.076 1.00 40.86 39.870 8.880 76.684 1.00 38.15 40.026 7.745 77.576 1.00 36.49 40.194 6.471 76.745 1.00 36.49 40.194 6.471 76.745 1.00 36.49 40.194 6.471 76.745 1.00 36.49 40.194 6.471 76.745 1.00 36.46 40.426 5.334 77.554 1.00 42.23 41.245 7.990 78.419 1.00 35.56	AAAA ATOM 2115 ND1 HIS AAAA ATOM 2117 CE1 HIS AAAA ATOM 2118 NE2 HIS AAAA ATOM 2120 C HIS AAAA ATOM 2121 O HIS AAAA ATOM 2122 N TYR AAAA ATOM 2124 CA TYR AAAA ATOM 2125 CB TYR AAAA ATOM 2126 CG TYR AAAA ATOM 2127 CD1 TYR AAAA ATOM 2128 CE1 TYR AAAA ATOM 2129 CD2 TYR AAAA ATOM 2130 CE2 TYR AAAA ATOM 2131 CZ TYR AAAA ATOM 2131 CZ TYR AAAA ATOM 2131 CZ TYR AAAA ATOM 2132 OH TYR AAAA ATOM 2135 O TYR AAAA ATOM 2136 N TYR AAAA ATOM 2136 N TYR AAAA ATOM 2138 CA TYR AAAA ATOM 2139 CB TYR AAAA ATOM 2139 CB TYR AAAA ATOM 2139 CB TYR AAAA ATOM 2140 CG TYR AAAA ATOM 2141 CD1 TYR AAAA ATOM 2141 CD1 TYR AAAA ATOM 2142 CE1 TYR AAAA ATOM 2144 CE2 TYR AAAA ATOM 2144 CE2 TYR AAAA ATOM 2144 CE2 TYR AAAA ATOM 2145 CZ TYR AAAA ATOM 2146 OH TYR AAAA ATOM 2148 C TYR AAAA ATOM 2150 N TYR AAAA ATOM 2150 N TYR AAAA ATOM 2150 N TYR AAAA ATOM 2150 CB TYR AAAA ATOM 2155 CD1 TYR AAAA ATOM 2155 CD1 TYR AAAA ATOM 2155 CD1 TYR AAAA ATOM 2156 CE1 TYR	223       39.432       23.726       78.115       1.00       43.35         223       38.242       23.594       78.666       1.00       44.77         223       37.911       22.314       78.640       1.00       45.50         223       42.529       20.994       73.969       1.00       41.67         224       43.386       21.939       75.843       1.00       40.87         224       44.740       22.109       75.351       1.00       39.54         224       44.490       24.297       74.345       1.00       43.98         224       44.499       24.297       74.345       1.00       43.98         224       44.499       24.297       74.345       1.00       45.59         224       43.411       25.121       74.561       1.00       42.61         224       42.771       25.714       73.517       1.00       45.59         224       44.310       24.671       71.988       1.00       47.82         224       43.218       25.491       72.231       1.00       48.72         224       45.804       21.284       76.000       1.00       39.39	ANAX ANAX ANAX ANAX ANAX ANAX ANAX ANAX
ATCH 2013 CB PRO 212 (0.986 6.909 8).999 1.00 48.56 AAAA ATCH 2181 C VAL 229 48.513 10.16 77.771 1.00 17.11 AAAA ATCH 2181 C VAL 229 48.513 10.16 77.771 1.00 17.11 AAAA ATCH 2181 C VAL 229 48.513 11.65 77.771 1.00 17.11 AAAA ATCH 2187 C VAL 229 48.513 11.65 77.771 1.00 17.11 AAAA ATCH 2187 C VAL 229 48.513 11.65 77.771 1.00 17.11 AAAA ATCH 2187 C VAL 229 48.513 11.65 77.771 1.00 17.11 AAAA ATCH 2187 C VAL 229 48.513 1.65 77.771 1.00 17.11 AAAA ATCH 2187 C VAL 229 48.513 1.65 77.771 1.00 17.11 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05 77.14 1.00 17.14 AAAA ATCH 2187 C VAL 229 18.05	ATOM 1997 CA CYS 209 ATOM 1998 C CYS 209 ATOM 1999 O CYS 209 ATOM 2000 CB CYS 209 ATOM 2001 SG CYS 209 ATOM 2002 N SER 210 ATOM 2004 CA SER 210 ATOM 2005 CB SER 210 ATOM 2006 OG SER 210 ATOM 2008 C SER 210 ATOM 2009 O SER 210 ATOM 2010 N ALA 211 ATOM 2012 CA ALA 211 ATOM 2013 CB ALA 211 ATOM 2014 C ALA 211 ATOM 2015 O ALA 211 ATOM 2016 N PRO 212 ATOM 2017 CD PRO 212	43.029       7.065       79.809       1.00       43.02         43.688       5.783       80.251       1.00       44.54         43.236       4.682       79.939       1.00       45.60         42.699       7.837       81.063       1.00       43.03         40.927       7.852       81.449       1.00       52.23         44.760       5.961       81.020       1.00       46.33         45.553       4.858       81.565       1.00       46.93         47.039       5.091       81.272       1.00       45.65         47.443       6.393       81.659       1.00       47.68         45.354       4.737       83.079       1.00       46.56         45.485       3.656       83.660       1.00       46.51         45.043       5.864       83.705       1.00       46.73         46.141       6.241       85.848       1.00       48.68         43.783       6.964       85.472       1.00       47.65         43.684       8.016       84.824       1.00       49.87         42.983       6.701       86.500       1.00       46.58         42.993	AAAA ATOM 2158 CE2 TYR AAAA ATOM 2159 CZ TYR AAAA ATOM 2160 OH TYR AAAA ATOM 2162 C TYR AAAA ATOM 2163 O TYR AAAA ATOM 2164 N ALA AAAA ATOM 2166 CA ALA AAAA ATOM 2166 CA ALA AAAA ATOM 2167 CB ALA AAAA ATOM 2168 C ALA AAAA ATOM 2169 O ALA AAAA ATOM 2170 N GLY AAAA ATOM 2171 CA GLY AAAA ATOM 2171 C GLY AAAA ATOM 2171 CA VAL AAAA ATOM 2177 CA VAL AAAA ATOM 2178 CB VAL AAAA ATOM 2178 CB VAL AAAA ATOM 2179 CG1 VAL	226       53.365       24.706       77.607       1.00 64.51         226       53.882       24.489       78.876       1.00 65.38         226       55.030       25.136       79.280       1.00 68.79         226       51.725       20.315       78.003       1.00 41.20         226       51.392       20.007       79.141       1.00 42.16         227       52.931       20.069       77.521       1.00 39.70         227       53.933       19.456       78.350       1.00 36.97         227       53.403       18.219       79.019       1.00 38.99         227       53.403       18.219       79.019       1.00 37.90         227       53.181       18.206       80.214       1.00 40.28         228       53.219       17.160       78.265       1.00 38.53         228       52.708       15.952       78.873       1.00 41.07         228       52.708       15.952       78.873       1.00 41.07         228       50.300       17.310       79.602       1.00 41.21         229       49.619       17.497       80.414       1.00 38.16         229       49.981       18.276       81	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA
ATOM 2083 C ALA 220 43.049 13.954 75.408 1.00 33.38 AAAA ATOM 2241 C ASN 237 53.972 31.320 76.872 1.00 37.18 AAAA	ATOM 2019 CB PRO 212 ATOM 2020 CG PRO 212 ATOM 2021 C PRO 212 ATOM 2021 C PRO 212 ATOM 2023 N ASP 213 ATOM 2025 CA ASP 213 ATOM 2026 CB ASP 213 ATOM 2027 C ASP 213 ATOM 2027 C ASP 213 ATOM 2028 O ASP 213 ATOM 2029 N ASN 214 ATOM 2031 CA ASN 214 ATOM 2031 CA ASN 214 ATOM 2031 CA ASN 214 ATOM 2033 CG ASN 214 ATOM 2033 CG ASN 214 ATOM 2034 OD1 ASN 214 ATOM 2035 ND2 ASN 214 ATOM 2038 C ASN 214 ATOM 2039 O ASN 214 ATOM 2040 N ASP 215 ATOM 2040 CA ASP 215 ATOM 2041 CB ASP 215 ATOM 2042 CA ASP 215 ATOM 2044 CG ASP 215 ATOM 2044 CG ASP 215 ATOM 2045 OD1 ASP 215 ATOM 2046 OD2 ASP 215 ATOM 2047 C ASP 215 ATOM 2048 O ASP 215 ATOM 2048 C ASP 215 ATOM 2048 C ASP 215 ATOM 2051 CA THR 216 ATOM 2052 CB THR 216 ATOM 2055 CG2 THR 216 ATOM 2055 CG2 THR 216 ATOM 2056 C THR 216 ATOM 2056 C THR 216 ATOM 2057 O THR 216 ATOM 2058 N ALA 217 ATOM 2060 CA ALA 217 ATOM 2060 CA ALA 217 ATOM 2061 CB ALA 217 ATOM 2066 CA CYS 218 ATOM 2067 C CYS 218 ATOM 2069 CB CYS 218 ATOM 2071 N VAL 219 ATOM 2076 CG2 VAL 219 ATOM 2077 C VAL 219 ATOM 2078 O VAL 219 ATOM 2079 N ALA 220 ATOM 2082 CB ALA 220	40.986 6.909 87.599 1.00 48.56 41.818 5.827 88.328 1.00 47.92 42.526 8.922 87.549 1.00 47.05 43.716 9.019 87.832 1.00 46.13 41.632 9.859 87.827 1.00 49.15 41.954 11.096 88.514 1.00 52.16 41.777 10.894 90.011 1.00 53.04 43.330 11.696 88.226 1.00 53.72 43.800 12.553 88.988 1.00 56.74 43.954 11.279 87.121 1.00 54.89 45.293 11.763 86.732 1.00 52.97 46.270 10.586 86.735 1.00 52.97 47.684 11.020 86.517 1.00 58.70 47.939 12.191 86.269 1.00 63.23 48.620 10.082 86.603 1.00 61.43 45.314 12.454 85.356 1.00 49.03 45.083 11.806 84.334 1.00 48.71 45.615 13.753 85.343 1.00 44.94 45.631 14.549 84.111 1.00 42.89 45.490 16.038 84.420 1.00 40.15 46.648 16.571 85.228 1.00 42.93 46.540 16.38 84.420 1.00 40.15 46.648 16.571 85.228 1.00 42.93 47.689 15.885 85.297 1.00 46.15 46.842 14.369 83.215 1.00 42.53 47.094 15.178 82.319 1.00 44.17 47.586 13.303 83.440 1.00 41.36 48.755 13.042 82.636 1.00 38.69 50.029 13.142 83.482 1.00 39.62 50.078 12.030 84.391 1.00 40.10 50.059 14.466 84.274 1.00 34.60 48.607 11.632 82.110 1.00 42.63 47.154 9.662 81.968 1.00 42.02 46.714 8.836 83.168 1.00 42.04 47.154 9.662 81.968 1.00 42.02 46.714 8.836 83.168 1.00 42.02 44.734 10.902 79.212 1.00 40.07 45.141 10.160 77.970 1.00 39.42 44.551 12.356 78.822 1.00 19.90 44.224 13.387 80.287 1.00 43.13 45.580 8.517 80.287 1.00 42.66 47.458 11.026 82.408 1.00 42.66 47.458 11.026 79.212 1.00 40.07 44.734 10.902 79.212 1.00 40.07 45.141 10.160 77.970 1.00 39.42 44.734 10.902 79.212 1.00 40.07 45.141 10.160 77.970 1.00 39.42 44.551 12.356 78.822 1.00 19.90 44.224 13.387 80.287 1.00 47.33 44.593 10.195 76.960 1.00 32.25 44.631 9.543 75.723 1.00 30.09 44.224 13.387 80.269 1.00 47.33 44.993 10.195 76.960 1.00 32.25 44.661 9.549 77.921 1.00 40.07 45.141 10.160 67.4745 1.00 30.065 44.221 7.646 74.212 1.00 32.25 44.381 1.0666 74.745 1.00 30.065 44.224 13.387 75.249 1.00 29.960 44.3861 2.531 73.397 1.00 29.93	AAAA ATOM 2181 C VAL AAAA ATOM 2182 O VAL AAAA ATOM 2183 N CYS AAAA ATOM 2185 CA CYS AAAA ATOM 2186 C CYS AAAA ATOM 2186 C CYS AAAA ATOM 2187 O CYS AAAA ATOM 2188 CB CYS AAAA ATOM 2189 SG CYS AAAA ATOM 2189 SG CYS AAAA ATOM 2190 N VAL AAAA ATOM 2191 CA VAL AAAA ATOM 2191 CG1 VAL AAAA ATOM 2191 CG1 VAL AAAA ATOM 2195 CG2 VAL AAAA ATOM 2195 CG2 VAL AAAA ATOM 2196 C VAL AAAA ATOM 2197 O VAL AAAA ATOM 2198 N PRO AAAA ATOM 2199 CD PRO AAAA ATOM 2199 CD PRO AAAA ATOM 2200 CA PRO AAAA ATOM 2201 CB PRO AAAA ATOM 2202 CG PRO AAAA ATOM 2208 CB ALA AAAA ATOM 2207 CA ALA AAAA ATOM 2208 CB ALA AAAA ATOM 2208 CB ALA AAAA ATOM 2208 CB ALA AAAA ATOM 2209 C ALA AAAA ATOM 2200 CA PRO AAAA ATOM 2211 N CYS AAAA ATOM 2211 CB PRO AAAA ATOM 2211 CB PRO AAAA ATOM 2212 CB PRO AAAA ATOM 2212 CB PRO AAAA ATOM 2213 CB PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2222 CG PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2222 CG PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2222 CG PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2222 CG PRO AAAA ATOM 2221 CB PRO AAAA ATOM 2222 CG PRO AAAA ATOM 2223 C PRO AAAA ATOM 2223 C PRO AAAA ATOM 2223 C PRO AAAA ATOM 2223 CB PRO AAAA ATOM 2223 CB PRO AAAA ATOM 2234 CA ASN AAAA ATOM 2234 CA ASN AAAA ATOM 2235 CB ASN AAAA ATOM 2236 CB ASN AAAA ATOM 2237 ODI ASN AAAA ATOM 2238 ND2 ASN	229       48.418       18.140       79.771       1.00       37.21         229       48.523       19.169       79.113       1.00       36.96         230       47.260       17.537       80.000       1.00       34.76         230       45.951       19.234       80.278       1.00       32.61         230       45.075       19.073       81.390       1.00       30.21         230       44.975       16.968       79.427       1.00       34.04         230       45.231       15.838       78.035       1.00       32.96         231       45.832       20.411       79.705       1.00       32.99         231       45.562       21.716       80.294       1.00       35.02         231       46.654       22.713       79.847       1.00       32.73         231       46.654       22.713       79.847       1.00       32.73         231       44.621       23.964       80.671       1.00       32.73         231       44.504       22.289       79.897       1.00       39.89         231       43.737       22.147       78.771       1.00       40.63	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA

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ATOM A  A  A  A  A  A  A  A  A  A  A  A  A	2268 2269 2270 2271 2272 2274	CC TYR CD1 TYR CE1 TYR CD2 TYR CE2 TYR CZ TYR OH TYR C TYR O TYR O TYR O ARG CA	239 239 239 239 239 239 239 240 240 240 240 240 240 240	52.375 53.015 50.304 50.927 52.272 52.854 49.166 49.793 47.852 47.110 45.626 45.255 43.771 43.117 43.256	31.957 33.016 33.070 34.119 34.094 35.161 28.660 27.662 28.768 27.655 27.940 29.132 29.132 29.143 30.082 31.392 31.888	72.680 73.260 72.938 73.515 73.679 74.301 72.416 72.091 72.201 71.583 71.580 72.421 72.648 71.753 71.869 72.842	1.00 27.41 1.00 26.72 1.00 26.63 1.00 27.10 1.00 27.10 1.00 32.09 1.00 32.09 1.00 34.50 1.00 30.65 1.00 27.19 1.00 25.69 1.00 26.86 1.00 29.42 1.00 30.93 1.00 33.99	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA	MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2420 CA ASN 2421 CB ASN 2422 CG ASN 2423 OD1 ASN 2424 ND2 ASN 2427 C ASN 2428 O ASN 2429 N ILE 2431 CA ILE 2432 CB ILE 2433 CG2 ILE 2434 CG1 ILE 2435 CD1 ILE 2436 C ILE 2437 O ILE 2438 N LEU	254 254 254 254 254 255 255 255 255 255	62.825 63.493 62.817 59.966 60.551 58.718 57.827 56.375 55.585 55.740 54.400 58.126 58.351 58.143	26.739 28.053 27.050 27.514 27.423 26.688 27.238 26.778 25.631 27.819	64.508 64.764 63.846 66.008 63.261 63.150 62.854 62.285 62.515 61.248 63.625 64.112 60.787 60.393 59.960	1.00 41.51 1.00 42.41 1.00 45.51 1.00 45.81 1.00 45.62 1.00 47.11 1.00 44.95 1.00 43.91 1.00 32.00 1.00 46.54 1.00 48.08 1.00 48.08	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	228356789991234567899012345678901234568901234567892323333333333333333333333333333333333	C ARG O ARG N PHE CA PHE CB PHE CG PHE CD1 PHE CD2 PHE CE1 PHE CE2 PHE C	22222211111112222222222222222222222222	47.6038 47.6038 47.5344 48.5344 49.52146 50.626	27.535 28.509 26.357 26.040 26.654 26.569 24.367 26.989 24.989 23.989 24.989 23.768 23.1916 23.768 23.1916 23.768 24.1916 22.741 23.768 24.1916 22.741 23.751 23	70.168 69.603 69.583 68.298 66.966.753 66.753 66.753 66.753 66.768 66.768 66.768 66.768 66.768 66.768 66.768 66.768 66.768 67.978 68.76	1.00 23.37 1.00 26.68 1.00 20.36 1.00 20.36 1.00 19.13 1.00 20.49 1.00 15.08 1.00 16.72 1.00 20.68 1.00 20.21 1.00 21.39 1.00 21.32 1.00 25.78 1.00 25.33 1.00 25.74 1.00 28.23 1.00 25.74 1.00 28.23 1.00 27.15 1.00 27.68 1.00 27.15 1.00 27.15 1.00 27.68 1.00 27.71 1.00 27.68 1.00 27.68 1.00 27.71 1.00 27.68 1.00 27.68 1.00 27.68 1.00 27.68 1.00 27.68 1.00 27.68 1.00 27.68 1.00 27.68 1.00 27.68 1.00 27.63 1.00 27.63 1.00 27.63 1.00 27.63 1.00 27.89 1.00 27.92	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2441 CB LEU 2442 CG LEU 2443 CD1 LEU 2444 CD2 LEU 2444 CD2 LEU 2445 C LEU 2446 O LEU 2447 N SER 2449 CA SER 2451 OG SER 2451 OG SER 2451 CA ALA 2458 CB ALA 2457 CA ALA 2458 CB ALA 2459 C ALA 2460 O ALA 2461 N GLU 2463 CA GLU 2466 CD GLU 2466 CD GLU 2467 OE1 GLU 2468 CC GLU 2467 OE1 GLU 2468 CC GLU 2471 N SER 2479 CA ASP 2481 CA SER 2482 CB SER 2483 OG SER 2484 CA SER 2484 CA SER 2484 CA SER 2489 CA ASP 2491 CB ASP 2491 CB ASP 2491 CB ASP 2492 CB SER 2493 CB SER 2495 CB SER 2496 CB SER 2497 CB SER 2498 CA SER 2499 CB SER	256 256 256 256 256 257 257 257 257 257 257 258 258 258	56.322 56.61213 56.62.755 60.7155 60.7155 62.750148 62.750148 62.951378 62.951378 62.951378 62.951378 62.951378 62.951378 62.951378 62.951378 62.951378 62.951378 62.951378 62.951378 63.8551378 63.8551378 63.8551378 63.8551378 63.8551378 63.85513 63.85	26.206 26.708 26.445 25.233 25.512 26.472 24.779 28.177 29.058 28.409 29.733 29.606 28.470 30.448 30.494 31.013 31.704 31.259 33.878 33.878 33.878 35.213 35.213 35.264 35.564 35.564 35.564 35.564 35.564 35.770 36.985 34.876 35.292	57.7155 57.2639 57.2639 57.2639 57.2639 57.3916 57.	1.00 48.70 1.00 49.20 1.00 49.20 1.00 46.78 1.00 49.57 1.00 54.86 1.00 55.37 1.00 57.82 1.00 61.01 1.00 63.24 1.00 65.73 1.00 68.76 1.00 69.08 1.00 70.99 1.00 74.18 1.00 70.65 1.00 68.39 1.00 70.62 1.00 74.23 1.00 70.62 1.00 74.23 1.00 76.56 1.00 66.31 1.00 66.31 1.00 66.31 1.00 66.31 1.00 66.89 1.00 66.89 1.00 66.89 1.00 57.33 1.00 57.08 1.00 57.08 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33 1.00 57.33	**************************************
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	12334467890234567801234567901235692346789012349012349012349012333333333333333333333333333333333333	NH2 ARG GYSS ARG GYSS SYSS ON CA ASP PHEE CYYSS ALA ALA ALA ASP PHEE CYYSS ALA ALA ALA ALA ALA ALA ALA ALA ALA A	22222222222222222222222222222222222222	44450120994576834834094226375894575559837744028100455555555555555555555555555555555555	23.940 24.847 24.847 25.341 25.361 25.361 25.361 25.361 25.361 25.361 25.361 25.361 26.5281 27.912 28.857 27.912 28.857 27.912 28.857 27.912 28.857 28.85	77777777777777777777777777777777777777	1.00 33.97 1.00 32.74 1.00 37.31 1.00 35.64 1.00 34.05 1.00 34.18 1.00 37.10 1.00 36.37 1.00 37.03 1.00 31.93 1.00 28.32 1.00 25.06 1.00 22.57 1.00 18.15 1.00 31.56 1.00 32.10 1.00 37.82 1.00 39.38 1.00 44.56 1.00 45.44 1.00 49.76 1.00 39.38 1.00 34.73 1.00 39.76 1.00 39.64 1.00 34.73 1.00 29.21 1.00 35.64 1.00 36.09 1.00 36.09 1.00 36.69 1.00 36.69 1.00 36.69 1.00 36.69 1.00 36.71 1.00 39.94	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2500 CG SER 2502 CG SER 2503 O SER 2503 CG SER 2504 CA GUU 2506 CG GUU 2507 CG GUU 2508 CG GUU 2510 OE2 2511 CA GUY 2512 C GUY 2513 CA GUY 2514 CA GUY 2515 CA CD1 PHE 2516 CA CD2 PHE 2521 CB PHE 2522 CG PHE 2523 CC PHE 2524 CD2 PHE 2525 CC CE2 2528 C CD2 PHE 2526 CC CE1 2531 CA LLE 2533 CA LLE 2534 CB LLE 2535 CG LLE 2536 CG LLE 2536 CG LLE 2547 CA LLE 2548 CA LLE 2547 CA LLE 2548 CA LLE 2547 CA LLE 2548 CA LLE 2546 CA LLE 2547 CA LLE 2546 CA LLE 2547 CA LLE 2546 CA LLE 2547 CA LLE 2546 CA LLE 2546 CA LLE 2547 CA LLE 2547 CA LLE 2547 CA CLE 2557 CA CLE 2577 CA GLY 2577 CA GLY 2577 CA GLY 2577 CA CLE	22222222222222222222222222222222222222	2536449682257648405736616375246802772272357716417456535555555555555555555555555555555555	35.66.7356.7356.7356.7356.7351.335.333.333.333.333.3333.3333333333	955555555555555555566656666556666666666	1.00 39.52 1.00 37.28 1.00 38.87 1.00 34.72 1.00 32.04 1.00 28.12 1.00 28.03 1.00 32.68 1.00 32.68 1.00 32.68 1.00 31.90 1.00 33.49 1.00 34.38 1.00 29.24 1.00 24.65 1.00 17.09 1.00 17.09 1.00 17.09 1.00 17.09 1.00 17.09 1.00 25.36 1.00 27.40 1.00 25.36 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.40 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90 1.00 27.94 1.00 31.90	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA

ATOM 2578 CB GLU 272 ATOM 2579 CG GLU 272    2580 CD GLU 272   2581 OE1 GLU 272 ATOM 2583 C GLU 272 ATOM 2583 C GLU 272 ATOM 2584 O GLU 272 ATOM 2585 N CYS 273 ATOM 2588 C CYS 273 ATOM 2588 C CYS 273 ATOM 2589 O CYS 273 ATOM 2590 CB CYS 273 ATOM 2591 SG CYS 273 ATOM 2591 SG CYS 273 ATOM 2592 N MET 274 ATOM 2595 CB MET 274 ATOM 2598 C MET 274 ATOM 2598 C MET 274 ATOM 2599 C MET 274 ATOM 2599 C MET 274 ATOM 2599 C MET 274 ATOM 2598 C MET 274 ATOM 2599 C MET 274 ATOM 2599 C MET 274 ATOM 2601 N GLN 275 ATOM 2603 CA GLN 275 ATOM 2604 CB GLN 275 ATOM 2605 CG GLN 275 ATOM 2605 CG GLN 275 ATOM 2606 CD GLN 275 ATOM 2607 OE1 GLN 275 ATOM 2608 NEZ GLN 275 ATOM 2611 C GLN 275 ATOM 2612 O GLN 275 ATOM 2615 CA GLU 276 ATOM 2616 CB GLU 276 ATOM 2617 CG GLU 276 ATOM 2618 CD GLU 276 ATOM 2618 CD GLU 276 ATOM 2619 OE1 GLU 276 ATOM 2620 OE2 GLU 276 ATOM 2621 C GLU 276 ATOM 2622 O CYS 277 ATOM 2623 CA CYS 277 ATOM 2624 C GYS 277 ATOM 2625 CA CYS 277 ATOM 2626 C CYS 277 ATOM 2627 C CYS 277 ATOM 2628 CB CYS 277 ATOM 2629 CB CYS 277 ATOM 2630 N PRO 278 ATOM 2631 CD PRO 278 ATOM 2631 CD PRO 278 ATOM 2633 CA PRO 278 ATOM 2634 CG PRO 278 ATOM 2639 CA SER 279 ATOM 2630 N PRO 278 ATOM 2631 CD PRO 278 ATOM 2631 CD PRO 278 ATOM 2632 CA PRO 278 ATOM 2633 CB PRO 278 ATOM 2634 CG PRO 278 ATOM 2635 C PRO 278 ATOM 2636 O PRO 278 ATOM 2637 N SER 279 ATOM 2644 CB SER 279 ATOM 2645 C CYS 277 ATOM 2646 C CYS 277 ATOM 2646 C CYS 277 ATOM 2647 CA GLY 280 ATOM 2648 C CYS 279 ATOM 2648 C GLY 280 ATOM 2649 O GLY 280 ATOM 2649 O GLY 280 ATOM 2640 C B SER 279 ATOM 2641 CC G CYS 277 ATOM 2642 C C PRO 288 ATOM 2653 C PRO 288 ATOM 2654 C C PRO 288 ATOM 2655 C PHE 281 ATOM 2650 N PHE 281	45.821 32.211 68.965 1.00 27.11 45.843 31.463 67.702 1.00 33.51 44.495 30.900 67.390 1.00 37.10 43.512 31.466 67.935 1.00 40.77 48.426 29.906 66.617 1.00 34.64 48.179 32.461 68.435 1.00 26.58 48.932 31.388 68.227 1.00 27.05 49.923 31.301 67.165 1.00 27.05 49.923 31.301 67.165 1.00 27.05 49.923 31.301 67.165 1.00 27.05 49.923 31.301 67.165 1.00 27.40 51.022 30.284 67.516 1.00 27.40 51.022 30.284 67.516 1.00 27.40 51.022 30.284 67.516 1.00 27.40 51.023 30.284 67.516 1.00 27.40 51.023 30.284 67.516 1.00 27.40 51.023 30.284 67.516 1.00 26.84 49.187 31.744 64.914 1.00 29.52 48.562 31.487 63.637 1.00 28.78 47.160 32.129 63.586 1.00 30.44 47.079 31.654 62.873 1.00 28.78 47.163 32.084 62.873 1.00 29.64 50.165 33.038 62.815 1.00 31.59 49.416 31.506 61.387 1.00 32.42 50.195 31.980 60.257 1.00 30.77 50.110 30.968 59.109 1.00 36.00 49.036 31.270 58.072 1.00 40.05 49.158 30.416 56.816 1.00 43.67 49.623 29.266 56.864 1.00 43.67 49.623 29.266 56.864 1.00 43.67 49.633 33.79 60.000 1.00 27.95 50.585 33.969 59.041 1.00 30.85 48.803 33.779 60.000 1.00 27.95 50.585 33.969 59.041 1.00 30.85 48.803 33.779 60.000 1.00 27.95 50.586 33.969 59.041 1.00 30.85 48.801 34.898 56.183 1.00 49.14 48.924 36.124 56.452 1.00 49.17 49.400 34.345 55.271 1.00 40.78 47.352 35.897 60.499 1.00 28.03 48.737 30.978 55.676 1.00 47.81 48.801 34.898 56.183 1.00 31.66 47.513 37.209 60.488 1.00 28.03 46.673 38.035 61.499 1.00 28.03 46.673 39.492 61.173 1.00 30.57 49.158 30.416 56.452 1.00 49.17 49.400 34.345 55.271 1.00 60.78 47.513 37.209 60.488 1.00 37.70 42.412 38.332 63.941 1.00 37.99 43.011 38.077 61.908 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 37.70 42.412 38.332 63.943 1.00 42.41 48.594 34.106 99.50 1.00 42.13 40.358 41.169 60.925 1.00 44.59 41.272 42.263 59.197 1.00 42.13 40.358 41.169 60.925 1.00 44.59	AAAA ATOM 2739 O MET 289 AAAA ATOM 2740 N TYR 290 AAAA ATOM 2741 CB TYR 290 AAAA ATOM 2743 CB TYR 290 AAAA ATOM 2744 CG TYR 290 AAAA ATOM 2745 CDI TYR 290 AAAA ATOM 2746 CDI TYR 290 AAAA ATOM 2746 CDI TYR 290 AAAA ATOM 2746 CDI TYR 290 AAAA ATOM 2747 CDZ TYR 290 AAAA ATOM 2748 CCZ TYR 290 AAAA ATOM 2748 CCZ TYR 290 AAAA ATOM 2750 OH TYR 290 AAAA ATOM 2750 OH TYR 290 AAAA ATOM 2751 O TYR 290 AAAA ATOM 2751 O TYR 290 AAAA ATOM 2752 C TYR 290 AAAA ATOM 2753 O TYR 290 AAAA ATOM 2756 CA CYS 291 AAAA ATOM 2756 CA CYS 291 AAAA ATOM 2757 C CYS 291 AAAA ATOM 2757 C CYS 291 AAAA ATOM 2758 O CYS 291 AAAA ATOM 2760 SG CYS 291 AAAA ATOM 2760 SG CYS 291 AAAA ATOM 2761 N ILE 292 AAAA ATOM 2763 CA ILE 292 AAAA ATOM 2766 CGI ILE 292 AAAA ATOM 2766 CGI ILE 292 AAAA ATOM 2766 CGI ILE 292 AAAA ATOM 2768 C ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2768 C ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2768 C ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2767 CDI ILE 292 AAAA ATOM 2768 C ILE 292 AAAA ATOM 2768 C ILE 292 AAAA ATOM 2769 O ILE 292 AAAA ATOM 2770 N PRO 293 AAAA ATOM 2771 CD PRO 293 AAAA ATOM 2772 CA PRO 293 AAAA ATOM 2773 CB PRO 293 AAAA ATOM 2774 CG PRO 293 AAAA ATOM 2775 C PRO 293 AAAA ATOM 2779 CA CYS 294 AAAA ATOM 2779 CA CYS 294 AAAA ATOM 2779 CA CYS 294 AAAA ATOM 2780 C CYS 298 AAAA ATOM 2780 C D ROO 297 AAAA ATOM 2800 C PRO 297 AAAA ATOM 2800 C PRO 297 AAAA ATOM 2800 C	52.501 38.544 65.931 1.00 28.48 AAAA S3.445 40.597 65.744 1.00 28.07 AAAA S3.445 40.597 65.744 1.00 28.07 AAAA S2.989 42.558 67.114 1.00 28.08 AAAA S3.52.989 42.558 67.114 1.00 28.08 AAAA S3.57.00 42.094 8.358 1.00 28.56 AAAA S5.046 41.761 68.336 1.00 26.58 AAAA S5.046 41.761 68.336 1.00 26.68 AAAA S5.046 41.761 68.336 1.00 26.68 AAAA S5.046 41.761 68.336 1.00 21.77 AAAA S5.046 41.501 70.696 1.00 21.77 AAAA S5.047 64 1.196 70.644 1.00 28.09 AAAA S5.047 64 41.196 70.644 1.00 28.09 AAAA S5.047 64 41.196 70.644 1.00 28.09 AAAA S5.047 64 41.196 70.644 1.00 34.58 AAAA S5.047 64 41.196 70.644 1.00 34.58 AAAA S5.1.395 41.998 65.281 1.00 30.98 AAAA S5.1.395 41.998 65.281 1.00 30.98 AAAA AAAA A48.287 41.394 65.560 1.00 21.25 AAAAA A48.287 41.394 65.560 1.00 21.25 AAAAA A48.287 41.095 66.314 1.00 36.19 AAAA A48.287 41.025 64.623 1.00 31.41 AAAA A48.287 41.025 64.623 1.00 31.41 AAAA A48.287 41.025 64.623 1.00 34.25 AAAA A48.552 39.512 63.781 1.00 34.25 AAAA A48.552 39.512 63.781 1.00 34.25 AAAA A48.552 39.512 63.781 1.00 34.25 AAAA A48.337 46.955 64.760 1.00 36.17 AAAA A48.337 46.955 64.760 1.00 36.37 AAAA A49.314 46.976 65.887 1.00 31.01 AAAA A49.314 66.976 65.887 1.00 31.01 AAAA A49.316 66.314 67.00 67.00 67.00 A49.314 66.976 65.887 1.00 6
ATOM 2654 CG PHE 281  ATOM 2655 CD1 PHE 281  ATOM 2656 CD2 PHE 281  ATOM 2658 CE2 PHE 281  ATOM 2658 CE2 PHE 281  ATOM 2659 CZ PHE 281  ATOM 2660 C PHE 281  ATOM 2661 O PHE 281  ATOM 2661 O PHE 281  ATOM 2662 N ILE 282  ATOM 2664 CA ILE 282  ATOM 2665 CB ILE 282  ATOM 2666 CG2 ILE 282  ATOM 2666 CG2 ILE 282  ATOM 2666 CG2 ILE 282  ATOM 2667 CG1 ILE 282  ATOM 2669 C ILE 282  ATOM 2669 C ILE 282  ATOM 2669 C ILE 282  ATOM 2670 O ILE 282  ATOM 2671 N ARG 283  ATOM 2675 CG ARG 283  ATOM 2675 CG ARG 283  ATOM 2676 CD ARG 283  ATOM 2677 NE ARG 283  ATOM 2679 CZ ARG 283  ATOM 2680 NH1 ARG 283  ATOM 2680 NH2 ARG 283  ATOM 2688 N ASN 284  ATOM 2688 N ASN 284  ATOM 2689 C ARG 283  ATOM 2689 C B ASN 284  ATOM 2691 CB ASN 284  ATOM 2692 CG ASN 284  ATOM 2693 OD1 ASN 284  ATOM 2694 ND2 ASN 284  ATOM 2699 C ASN 284  ATOM 2698 N GLY 285  ATOM 2698 N GLY 285  ATOM 2700 CA GLY 285  ATOM 2701 C GLY 285  ATOM 2701 C GLY 285  ATOM 2700 CA SER 286  ATOM 2701 C GLY 285  ATOM 2702 C GLN 287  ATOM 2703 N SER 286  ATOM 2704 CB SER 286  ATOM 2705 CA SER 286  ATOM 2707 CG SER 286  ATOM 2708 C SER 286  ATOM 2709 C SER 286  ATOM 2701 C GLN 287  ATOM 2711 N GLN 287  ATOM 2712 C GLN 287  ATOM 2713 CA GLN 287  ATOM 2714 CB GLN 287  ATOM 2715 CG GLN 287  ATOM 2716 CD GLN 287  ATOM 2727 CG SER 288  ATOM 2731 N MET 289  ATOM 2731 N MET 289  ATOM 2732 C SER 288  ATOM 2733 CA MET 289  ATOM 2734 CB MET 289  ATOM 2736 CD MET 289  ATOM 2737 CE MET 289  ATOM 2738 C MET 289  ATOM 2739 C SER 288  ATOM 2731 C MET 289  ATOM 2731 C MET 289  ATOM 2733 C MET 289	44.431 43.565 64.761 1.00 40.38 42.536 42.345 63.728 1.00 36.40 41.828 42.663 64.794 1.00 37.95 42.321 43.440 65.827 1.00 40.52 45.688 43.114 60.312 1.00 19.30 45.688 42.211 59.475 1.00 15.84 46.648 44.011 60.414 1.00 40.76 47.821 44.030 59.562 1.00 40.08 47.823 45.367 58.750 1.00 14.68 48.695 46.000 56.504 1.00 33.54 48.695 46.000 56.504 1.00 35.73 48.951 43.966 60.604 1.00 42.12 48.811 44.584 61.667 1.00 45.18 50.037 43.220 60.366 1.00 37.15 52.564 42.717 59.933 1.00 37.15 52.565 42.059 61.290 1.00 33.34 53.162 40.325 60.001 1.00 32.46 52.204 39.347 59.531 1.00 33.43 53.760 38.076 58.480 1.00 41.70 51.768 44.514 61.416 1.00 13.81 52.706 44.514 60.371 1.00 33.81 53.760 38.076 58.480 1.00 41.70 51.768 44.514 61.416 1.00 31.85 52.594 38.291 58.809 1.00 33.81 52.700 46.190 62.895 1.00 34.98 51.750 37.466 78.411 1.00 31.09 51.768 44.514 61.416 1.00 34.98 51.593 46.657 65.243 1.00 35.24 52.392 46.587 64.329 1.00 35.24 52.392 46.587 64.329 1.00 36.36 54.180 45.884 62.617 1.00 45.18 54.180 45.884 62.617 1.00 37.38 54.180 45.884 62.617 1.00 37.38 55.956 46.019 60.910 1.00 25.52 56.658 44.767 58.956 1.00 31.81 55.592 43.676 60.552 1.00 36.36 54.673 45.182 63.409 1.00 31.64 55.592 43.676 60.552 1.00 35.43 55.952 44.694 62.637 1.00 35.24 55.953 43.676 60.552 1.00 35.43 55.954 46.270 61.409 1.00 31.64 55.956 48.377 58.840 1.00 41.70 56.627 44.719 60.20 1.00 25.52 56.658 44.767 58.956 1.00 24.57 56.830 43.577 58.840 1.00 24.57 56.830 43.577 58.840 1.00 24.57 56.830 43.577 58.840 1.00 24.57 56.809 41.037 58.959 1.00 23.45 56.720 42.682 59.359 1.00 35.47 57.552 43.964 56.871 1.00 22.40 58.720 42.682 59.359 1.00 35.47 57.570 40.152 61.057 1.00 23.25 55.7670 40.152 61.057 1.00 23.25 55.7670 40.152 61.005 1.00 28.11 56.692 37.490 59.859 1.00 28.12 55.719 38.676 60.230 1.00 28.12 55.729 42.977 64.473 1.00 28.15 55.729 42.977 64.473 1.00 28.53 55.7670 40.152 61.005 1.00 27.92	AAAA ATOM 2812 CB PRO 299 AAAA ATOM 2814 C PRO 299 AAAA ATOM 2815 O PRO 299 AAAA ATOM 2815 O PRO 299 AAAA ATOM 2815 O PRO 299 AAAA ATOM 2816 N LYS 300 AAAA ATOM 2819 CB LYS 300 AAAA ATOM 2819 CB LYS 300 AAAA ATOM 2820 CC LYS 300 AAAA ATOM 2821 CD LYS 300 AAAA ATOM 2822 CE LYS 300 AAAA ATOM 2822 CE LYS 300 AAAA ATOM 2822 CE LYS 300 AAAA ATOM 2823 NZ LYS 300 AAAA ATOM 2823 NZ LYS 300 AAAA ATOM 2828 O LYS 300 AAAA ATOM 2831 CA VAL 301 AAAA ATOM 2831 CB VAL 301 AAAA ATOM 2831 CC VAL 301 AAAA ATOM 2833 CG VAL 301 AAAA ATOM 2834 CG2 VAL 301 AAAA ATOM 2837 N CYS 302 AAAA ATOM 2839 CA CYS 302 AAAA ATOM 2839 CA CYS 302 AAAA ATOM 2839 CA CYS 302 AAAA ATOM 2840 C CYS 302 AAAA ATOM 2840 C CYS 302 AAAA ATOM 2841 O CYS 302 AAAA ATOM 2842 CB CYS 302 AAAA ATOM 2844 N GLU 303 AAAA ATOM 2846 CA GLU 303 AAAA ATOM 2848 CG GLU 303 AAAA ATOM 2846 CA GLU 303 AAAA ATOM 2846 CA GLU 303 AAAA ATOM 2851 OEZ GLU 303 AAAA ATOM 2851 OEZ GLU 303 AAAA ATOM 2850 OEI GLU 303 AAAA ATOM 2850 CE GLU 304 AAAA ATOM 2850 CE GLU 304 AAAA ATOM 2850 CE GLU 304 AAAA ATOM 2850 CE GLU 305 AAAA ATOM 2860 CE GLU 305 AAA	47, 377 48.182 54.734 1.00 35.28 AAAA 46.229 47.013 52.162 1.00 28.64 AAAA 46.456 45.980 52.780 1.00 29.83 AAAA 46.478 47.095 50.843 1.00 25.99 AAAA 46.478 45.963 49.972 1.00 28.84 AAAA 45.076 44.637 48.341 1.00 28.84 AAAA 44.045 43.550 48.220 1.00 31.17 AAAA 44.045 43.550 48.220 1.00 31.17 AAAA 44.045 41.953 48.220 1.00 32.43 AAAA 44.045 41.953 48.221 1.00 38.15 AAAA 44.600 41.953 48.6422 1.00 38.15 AAAA 46.285 47.372 48.046 1.00 36.31 AAAA 48.670 46.573 48.961 1.00 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3087   ND2 ASN   329   51.058   3097   ND2 ASN   329   51.058   3097   ND2 ASN   329   51.058   3099   C ASN   329   46.805   3092   N LEU   330   44.703   3095   CB LEU   330   43.786   3096   CG LEU   330   43.786   3096   CG LEU   330   44.703   3095   CB LEU   330   44.291   3100   C LEU   330   341.076   3099   CD LEU   330   341.076   3100   C LEU   331   41.673   3101   N LEU   331   41.673   3101   N LEU   331   41.673   3103   CA LEU   331   41.938   3105   CC LEU   331   41.364   3106   CD1 LEU   331   41.364   3109   CD2 LEU   331   41.938   3105   CC LEU   331   31.938   31.938   31.938   31.938   31.938   31.938   31.938   31.938   31.938   31.938   31.938   31.938	1.00 42.67 1.00 42.67 1.00 41.64 1.00 41.62 1.00 41.76 1.00 41.76 1.00 41.76 1.00 41.76 1.00 41.76 1.00 41.76 1.00 41.76 1.00 42.53 1.00 42.53 1.00 50.14 1.00 51.13 1.00 38.93 1.00 33.40 1.00 37.51 1.00 28.43 1.00 37.51 1.00 27.60 1.00 36.21 1.00 41.14 1.00 43.85 1.00 44.14 1.00 43.85 1.00 49.33 1.00 44.48 1.00 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44.769 39.879 29 44.769 39.879 20 42.771 20 39.890 20 39.491 20 39.49	309	2991 N MET 2993 CA MET 2994 CB MET 2995 CG MET 2996 SD MET 2997 CE MET 2998 C MET 2999 O MET 3000 N LEU 3002 CA LEU 3003 CB LEU 3005 CD1 LEU 3006 CD2 LEU 3007 C LEU 3008 O LEU 3009 N GLN 3011 CA GLN 3012 CB GLN 3012 CB GLN 3013 CG GLN 3014 CD GLN 3015 OE1 GLN 3015 OE1 GLN 3016 NE2 GLN 3017 C GLN 3018 C GLY 3020 O GLN 3021 N GLY 3021 N GLY 3022 C GLY 3023 CA GLY 3024 C GLY 3025 O GLY 3026 N CYS 3028 CA CYS 3029 C CYS 3029 C CYS 3030 O CYS 3031 CB CYS 3032 SG CYS 3031 N THR 3035 CA THR 3035 CA THR	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM
1000   1000	3068   CD LYS   327   55.468   4			1.00 42.67 1.00 41.64 1.00 41.62 1.00 41.76 1.00 41.14 1.00 41.60 1.00 42.53 1.00 45.37 1.00 50.14 1.00 36.14 1.00 36.14 1.00 32.23 1.00 37.51 1.00 27.87 1.00 27.87 1.00 27.60 1.00 41.14 1.00 49.31 1.00 49.31 1.00 49.31 1.00 49.31 1.00 49.31 1.00 49.31 1.00 44.77 1.00 49.37 1.00 44.77 1.00 45.82 1.00 48.10 1.00 48.10 1.00 48.10 1.00 48.10 1.00 46.68 1.00 50.62 1.00 53.04 1.00 65.53 1.00 65.53 1.00 65.53 1.00 65.53 1.00 65.53 1.00 65.53 1.00 65.297 1.00 53.19 1.00 47.38 1.00 65.53 1.00 65.53 1.00 65.297 1.00 53.19 1.00 65.53 1.00 65.297 1.00 59.24 1.00 65.297 1.00 59.24 1.00 65.297 1.00 59.24 1.00 65.297 1.00 59.24 1.00 65.297 1.00 59.24 1.00 66.24 1.00 78.28 1.00 66.24 1.00 78.28 1.00 66.24 1.00 59.24 1.00 66.24 1.00 59.24 1.00 66.24 1.00 59.24 1.00 65.16 1.00 59.24 1.00 66.24 1.00 66.24 1.00 59.24 1.00 66.24 1.00 67.29 1.00 67.29	746 4 64.006 34.994 1.00 42.67 1459 43.901 35.155 1.00 41.62 133 43.319 16.1659 1.00 41.76 130 41.972 36.731 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31.00 64.134 313 34.531 45.593 39.141 1.00 43.53 313 35.006 42.774 31.312 66.00 44.48 313 35.006 42.774 31.312 66.00 44.77 313 34.531 45.593 39.141 1.00 43.53 313 35.006 42.774 31.315 66.00 44.77 313 34.531 45.593 39.141 1.00 46.56 313 35.006 42.774 31.315 66.00 44.77 313 34.531 47.594 30.791 1.00 46.56 313 37.404 39.40 30.771 1.00 46.56 313 37.606 41.795 31.00 46.69 31.313 31.00 46.85 313 35.006 42.776 31.319 1.00 66.77 313 34.531 47.96 30.771 1.00 66.77 314 31.294 40.606 31.713 1.00 66.56 314 31.33 35.006 42.776 31.391 1.00 66.57 315 31.394 40.609 31.713 1.00 66.56 313 37.706 47.795 35.600 30.00 67.79 314 31.404 31.500 47.795 35.600 30.00 67.79 315 31.306 44.795 31.00 66.600 30.713 316 41.796 43.896 42.795 31.00 66.600 30.707 317 37.212 44.166 31.777 1.00 66.24 318 33.396 48.290 48.290 1.00 65.16 319 40.895 37.896 37.897 30.00 66.29 319 39.609 41.396 41.397 30.00 66.69 319 39.609 41.396 41.397 30.00 66.69 319 39.609 41.396 41.397 30.00 66.69 319 39.609 41.396 41.397 30.0	2007   CS   LYS   309

-	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3225 O LEU 3226 N GLU 3228 CA GLU 3229 CB GLU 3230 CG GLU 3231 CD GLU 3232 OE1 GLU 3232 OE2 GLU 3234 C GLU 3235 O GLU 3236 N ASN 3236 N ASN 3238 CA ASN 3239 CB ASN 3240 CG ASN 3241 OD1 ASN	345 345 345 345 345 345 345 345 345 346 346 346 346	32.200 50.121 32.626 50.715 31.708 51.871 31.979 52.524 30.679 52.858 29.782 51.979 30.537 53.996 32.708 49.749 33.257 50.085 32.160 48.552 32.208 47.529 30.852 46.771 30.989 45.244 31.208 44.748	40.528 1.00 40.2 40.719 1.00 38.3 41.973 1.00 41.3 42.326 1.00 41.3 43.650 1.00 48.8 44.378 1.00 63.9 44.887 1.00 62.1 43.130 1.00 43.0 44.166 1.00 45.5 42.968 1.00 45.7 44.022 1.00 48.3 44.058 1.00 51.5 44.244 1.00 55.3 45.365 1.00 54.8	9 AAAA 2 AAAA 4 AAAA 5 AAAA 1 AAAA 2 AAAA 1 AAAA 5 AAAA 1 AAAA 8 AAAA 9 AAAA	MOTA HOTA HOTA HOTA HOTA HOTA HOTA HOTA H	3381 NE ARG 3383 CZ ARG 3384 NH1 ARG 3387 NH2 ARG 3390 C ARG 3391 O ARG 3392 N HIS 3394 CA HIS 3395 CB HIS 3396 CG HIS 3396 CG HIS 3397 CD2 HIS 3398 ND1 HIS 3400 CE1 HIS 3401 NE2 HIS	361 42.2 361 38.4 361 38.6 362 37.4 362 36.9 362 37.8 362 37.8 362 38.0 362 38.6 362 39.3 362 38.9 362 38.9 362 38.9	51.005 51.302 36.49.769 36.53.782 21.52.579 55.54.33 52.53.518 55.52.285 43.52.621 38.53.793 55.51.704 13.52.296 57.53.565 56.53.064	23.684 23.544 23.982 25.126 25.252 24.479 23.172 23.172 21.388 21.361 20.383 20.346 24.917	1.00 41.80 1.00 30.62 1.00 34.58 1.00 32.24 1.00 34.64 1.00 33.74 1.00 33.42 1.00 35.37 1.00 33.84 1.00 32.38 1.00 29.52 1.00 36.04	AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3242 ND2 ASN 3246 O ASN 3247 N PHE 3249 CA PHE 3250 CB PHE 3251 CG PHE 3252 CD1 PHE 3253 CD2 PHE 3255 CE2 PHE 3256 CZ PHE 3256 CZ PHE 3257 C PHE 3258 O PHE 3258 O PHE 3258 O PHE 3258 O PHE 3261 CA MET 3262 CB MET 3263 CG MET 3264 SD MET 3263 CG MET 3264 SD MET 3265 CE MET 3266 C MET 3267 O MET 3268 N GLY 3270 CA GLY 3271 C GLY 3271 C GLY 3272 O GLY 3273 N LEU 3275 CA LEU 3276 CB LEU 3277 CG LEU 3278 CD1 LEU 3277 CG LEU 3278 CD1 LEU 3278 CD1 LEU 3278 CD1 LEU 3279 CD2 LEU 3278 CD1 LEU 3278 CD1 LEU 3278 CD1 LEU 3277 CG LEU 3278 CD1 LEU 3278 CD1 LEU 3278 CD1 LEU 3279 CD2 LEU 3280 C LEU 3281 O LEU 3281 O LEU 3282 N ILE 3286 CG2 ILE 3287 CG1 ILE 3288 CD1 ILE 3289 C ILE 3299 C GLU 3291 N GLU 3293 CA GLU 3295 CG GLU 3297 OE1 GLU 3298 CE GLU 3299 C GLU	346 347 347 347 347 347 347 347 347	33.408 46.609  33.530 45.494  34.312 47.105  35.505 46.366  35.314 45.747  34.523 44.492  33.374 44.401  34.975 43.367  32.687 43.207  33.153 42.080  37.847 46.913  36.405 48.633  37.468 49.615  37.525 50.076  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.346 50.835  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.435 48.955  37.441 50.835  38.237 51.679  37.435 48.955  37.441 53.501  38.226 50.150  38.226 50.150  38.226 50.150  38.226 50.150  38.237 51.679  31.211 53.501  31.237  31.211 53.501  31.237  31.238  31.239  31.239  31.239  31.2304  31.237  31.237  31.238  31.237  31.238  31.237  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238  31.237  31.238	43.141 1.00 55.3  43.719 1.00 49.7  44.226 1.00 52.9  42.884 1.00 49.9  42.500 1.00 47.8  41.131 1.00 46.2  40.350 1.00 47.1  41.783 1.00 48.3  40.225 1.00 48.1  41.669 1.00 48.7  40.885 1.00 50.0  42.397 1.00 49.3  42.438 1.00 52.1  42.258 1.00 43.6  42.092 1.00 43.6  42.092 1.00 43.6  42.971 1.00 45.3  43.667 1.00 45.3  43.667 1.00 45.3  43.667 1.00 45.7  44.504 1.00 47.3  45.571 1.00 46.7  46.038 1.00 47.3  46.982 1.00 47.3  47.817 1.00 46.7  49.189 1.00 50.7  49.823 1.00 47.3  47.817 1.00 46.7  49.189 1.00 50.7  49.823 1.00 49.7  50.078 1.00 51.0  46.568 1.00 48.8  47.433 1.00 51.9  48.834 1.00 47.3  49.823 1.00 49.7  50.078 1.00 50.7  49.823 1.00 49.7  50.078 1.00 50.7  49.823 1.00 49.7  49.823 1.00 49.7  50.078 1.00 50.7  49.823 1.00 49.7  50.078 1.00 50.7  49.823 1.00 49.7  50.078 1.00 36.4  41.829 1.00 36.4  42.975 1.00 36.4  43.279 1.00 36.4  44.800 1.00 44.8  47.433 1.00 51.9  45.135 1.00 41.1  44.780 1.00 44.8  45.834 1.00 37.1  46.563 1.00 44.8  50.252 1.00 49.2  45.440 1.00 49.8  50.252 1.00 49.2	5 AAAA 5 AAAA 6 AAAA 6 AAAA 7 AAAA 7 AAAA 1 AAAA 1 AAAA 1 AAAA 1 AAAA 2 AAAA 2 AAAA 6 AAAA 6 AAAA 6 AAAA 6 AAAA 6 AAAA 7 AAAA 8 AAAA	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3404 O HIS 3405 N SER 3407 CA SER 3408 CB SER 3409 OG SER 3411 C SER 3412 O SER 3413 N HIS 3415 CA HIS 3416 CB HIS 3417 CG HIS 3417 CG HIS 3418 CD2 HIS 3421 CE1 HIS 3421 CE1 HIS 3422 NE2 HIS 3421 C HIS 3422 NE2 HIS 3423 C HIS 3424 C HIS 3425 O HIS 3426 N ALA 3427 CB ALA 3428 CA ALA 3428 CA ALA 3429 CB ALA 3431 O ALA 3432 N LEU 3435 CB LEU 3436 CG LEU 3437 CD1 LEU 3436 CG LEU 3437 CD1 LEU 3438 CD2 LEU 3439 C LEU 3437 CD1 LEU 3438 CD2 LEU 3437 CD1 LEU 3438 CD2 LEU 3439 C LEU 3443 CA VAL 3444 CB VAL 3445 CG1 VAL 3446 CG2 VAL 3446 CG2 VAL 3447 C VAL 3446 CG2 VAL 3446 CG2 VAL 3446 CG1 VAL 3446 CG2 VAL 3446 CG1 VAL 3446 CG2 VAL 3446 CG2 VAL 3446 CG1 VAL 3446 CG2 VAL 3446 CG1 VAL 3446 CG2 VAL	363 35.2 363 34.3 363 34.9 363 35.6 363 33.0 363 32.7 364 32.2 364 30.9 364 32.1 364 32.1 364 32.2 364 32.1 364 32.1 365 364 32.1 366 32.3 365 29.2 365 29.2 365 30.0 365 29.2 365 30.0 365 29.2 365 30.0 365 29.2 367 29.2 367 367 368 31.7 366 32.3 366 32.3 366 32.3 366 32.3 367 29.9 367 29.9 367 29.9 367 29.9 367 29.9 367 29.9 367 29.9 367 29.9 367 29.9 368 31.5 369 31.6	53.527 53.891 53.891 54.291 54.265 54.3265 54.364 55.265 54.364 55.208 55.213 55.247 55.247 55.247 55.247 55.247 55.247 55.247 55.247 56.863 60.329 56.803 60.006 60.00	26.013 27.103 28.442 28.465 28.965 27.645 22.645 22.976 23.454 22.976 23.454 22.976 23.454 22.976 23.454 22.976 23.454 22.976 23.454	1.00 37.14 1.00 38.68 1.00 37.92 1.00 35.98 1.00 40.17 1.00 38.73 1.00 43.35 1.00 45.91 1.00 42.21 1.00 40.39 1.00 37.68 1.00 37.68 1.00 36.18 1.00 37.68 1.00 37.88 1.00 42.23 1.00 42.23 1.00 42.23 1.00 42.62 1.00 42.62 1.00 37.83 1.00 34.63 1.00 37.85 1.00 37.85 1.00 38.87 1.00 34.89 1.00 34.89 1.00 34.89 1.00 34.89 1.00 34.89 1.00 34.89 1.00 36.70 1.00 34.89 1.00 36.70 1.00 36.70 1.00 37.44 1.00 36.70 1.00 37.44 1.00 36.70 1.00 37.44 1.00 36.70 1.00 37.44 1.00 36.70 1.00 37.44 1.00 36.70 1.00 37.44	**************************************
	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3301 N VAL 3303 CA VAL 3304 CB VAL 3305 CG1 VAL 3306 CG2 VAL 3307 C VAL 3308 O VAL 33111 CA VAL 3312 CB VAL 3313 CG1 VAL 3314 CG2 VAL 3315 C VAL 3317 N THR 3319 CA THR 3320 CB THR 3321 OG1 THR 3322 CG2 THR 3322 CG2 THR 3323 CG2 THR 3323 CG2 THR 3323 CG2 TTR 3323 CG TYR 3324 C THR 3325 O THR 3326 N GLY 3327 CG TYR 3328 CA GLY 3329 C GLY 3330 O GLY 3331 N TYR 3333 CA TYR 3333 CA TYR 3334 CB TYR 3334 CB TYR 3335 CG TYR 3336 CD1 TYR 3337 CE1 TYR 3333 CA TYR 3334 CB TYR 3343 C TYR 3344 O TYR 3345 N VAL 3347 CA VAL 3347 CA VAL 3348 CB VAL 3349 CG1 VAL 3347 CA VAL 3348 CB VAL 3349 CG1 VAL 3347 CA VAL 3348 CB VAL 3349 CG1 VAL 3350 CG2 VAL 3351 C VAL 3355 CA LYS 3356 CB LYS 3357 CG LYS 3358 CD LYS 3359 CE LYS 3359 CE LYS 3359 CE LYS 3359 CE LYS 3357 CG LYS 3357	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	6.662 55.087 7.897 55.025 9.152 55.491 7.657 55.856 6.921 54.179 6.915 52.947 7.096 54.798 7.465 54.064 6.562 54.373 5.139 53.868 8.861 54.640 9.006 53.779 1.289 54.170 9.876 53.789 1.289 54.170 2.193 53.045 1.701 51.789 2.173 53.026 9.876 53.779 1.289 54.170 2.193 53.045 1.701 51.789 2.173 53.026 9.876 53.778 1.740 54.579 1.289 54.170 9.876 53.778 9.889 55.36 0.652 54.445 0.203 56.319 9.950 54.979 9.502 56.301 9.484 53.708 0.652 54.445 0.651 53.373 9.484 53.019 9.553 52.011 7.605 55.248 7.470 54.025 6.583 56.085 5.199 55.635 4.181 56.307 9.484 93.725 4.880 54.918 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 4.891 55.498 5.998 54.293 5.874 53.822 2.499 54.937 2.290 53.712 1.536 55.647 9.108 56.323 0.131 55.498 4.890 54.918	44.866 1.00 36.3 44.129 1.00 32.1 45.040 1.00 30.6 44.287 1.00 32.7 46.262 1.00 27.4 42.948 1.00 32.2 40.568 1.00 32.5 39.336 1.00 27.4 38.126 1.00 20.0 39.599 1.00 29.9 40.374 1.00 34.2 40.167 1.00 34.3 40.401 1.00 35.1 40.401 1.00 35.1 40.401 1.00 35.1 40.401 1.00 36.7 40.483 1.00 41.4 42.497 1.00 34.3 38.988 1.00 32.4 38.988 1.00 32.4 38.988 1.00 27.2 31.732 1.00 22.0 31.732 1.00 23.3 35.962 1.00 27.9 36.538 1.00 27.2 34.785 1.00 23.3 35.962 1.00 27.9 36.538 1.00 27.2 31.732 1.00 23.3 31.732 1.00 22.0 31.732 1.00 23.3 35.962 1.00 27.9 36.538 1.00 27.2 34.785 1.00 23.3 35.962 1.00 27.9 36.538 1.00 27.2 31.732 1.00 22.0 31.732 1.00 22.0 31.732 1.00 22.0 31.732 1.00 23.9 34.482 1.00 26.5 30.566 1.00 23.7 29.928 1.00 23.9 31.461 1.00 25.7 33.762 1.00 27.0 33.762 1.00 27.0 33.545 1.00 23.9 34.482 1.00 26.2 34.499 1.00 24.3 27.640 1.00 23.9 31.828 1.00 20.0 31.327 1.00 22.1 29.984 1.00 24.3 27.640 1.00 23.9 31.828 1.00 23.9 32.152 1.00 27.7 29.343 1.00 25.7 29.343 1.00 25.7 29.343 1.00 25.9 29.861 1.00 30.6 29.572 1.00 27.7 29.343 1.00 25.9 29.999 1.00 26.5 31.496 1.00 30.6 29.572 1.00 27.7 29.343 1.00 25.9 29.999 1.00 26.5 31.496 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.5710 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6 20.57110 1.00 30.6	2         AAAA           3         AAAA           4         AAAA           5         AAAA           6         AAAA           6         AAAA           7         AAAA           8         AAAA	ATOM 3 AT	1466	370       32.13         370       33.13         370       33.13         371       33.23         371       32.84         371       34.70         371       34.70         371       34.50         371       34.50         371       34.80         372       37.81         372       36.80         372       37.20         372       37.20         372       37.20         373       35.70         373       35.70         373       35.70         373       35.70         373       35.70         373       35.70         374       37.80         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373       33.60         373	3 59.350 59.350 59.350 59.350 50.367 56.960 56.960 56.9851 56.9857 56.9857 56.9857 56.9857 56.9857 56.9857 56.9857 56.9857 56.9857 56.8857 56.887 56.887 56.887 56.887 56.887 57.8867 57.8	39.755 13 39.904 33 30.904 33 30.1613 33 30.	1.00 41.56 1.00 39.49 1.00 39.26 1.00 37.46 1.00 35.50 1.00 38.29 1.00 38.48 1.00 41.45 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.08 1.00 41.40 1.00 41.40 1.00 49.64 1.00 49.64 1.00 49.64 1.00 49.64 1.00 49.66 1.00 49.66 1.00 49.66 1.00 49.66 1.00 38.97 1.00 49.66 1.00 38.97 1.00 49.66 1.00 38.97 1.00 49.66 1.00 38.97 1.00 49.66 1.00 38.97 1.00 49.66 1.00 38.97 1.00 49.66 1.00 39.91 1.00 39.91 1.00 39.91	**************************************

ATOM 3551 CD2 LEU 377  ATOM 3552 C LEU 377  3553 O LEU 377  3554 N ILE 378  AL 3556 CA ILE 378  ATOM 3557 CB ILE 378  ATOM 3558 CG2 ILE 378  ATOM 3559 CG1 ILE 378  ATOM 3560 CD1 ILE 378  ATOM 3561 C ILE 378  ATOM 3561 C ILE 378  ATOM 3563 N LEU 379  ATOM 3565 CA LEU 379  ATOM 3565 CA LEU 379  ATOM 3566 CB LEU 379  ATOM 3567 CG LEU 379  ATOM 3568 CD1 LEU 379  ATOM 3568 CD1 LEU 379  ATOM 3570 C LEU 379  ATOM 3571 O LEU 379  ATOM 3571 O LEU 379  ATOM 3572 N GLY 380  ATOM 3575 C GLY 380  ATOM 3576 O GLY 380  ATOM 3576 O GLY 380  ATOM 3577 N GLU 381  ATOM 3579 CA GLU 381  ATOM 3579 CA GLU 381	48.524 59.490 46.127 1.00 35.51 AA 46.966 59.205 42.276 1.00 35.54 AA 46.792 57.988 42.342 1.00 32.60 AA 47.603 59.800 41.267 1.00 35.90 AA 48.258 59.087 40.169 1.00 37.38 AA 47.621 59.424 38.797 1.00 36.92 AA 48.421 58.788 37.675 1.00 31.15 AA 46.186 58.878 38.737 1.00 36.11 AA 45.373 59.423 37.585 1.00 33.62 AA 49.727 59.543 40.194 1.00 39.52 AA 49.727 59.543 40.194 1.00 39.52 AA 50.032 60.736 40.099 1.00 40.21 AA 50.628 58.577 40.345 1.00 40.05 AA 52.055 58.842 40.447 1.00 38.95 AA 52.739 57.650 41.129 1.00 38.88 AA 52.095 57.212 42.463 1.00 38.74 AA 52.835 56.032 43.029 1.00 37.60 AA 52.835 56.032 43.029 1.00 37.60 AA 52.956 58.755 36.778 1.00 36.71 AA 52.956 58.755 36.778 1.00 34.15 AA 54.229 57.991 36.485 1.00 36.80 AA 54.705 57.186 37.432 1.00 36.39 AA 55.936 56.393 37.259 1.00 36.23 AA	ATOM 3707 C LEU 393 ATOM 3708 O LEU 393 ATOM 3708 O LEU 393 ATOM 3709 N ASP 394 ATOM 3711 CA ASP 394 ATOM 3712 CB ASP 394 ATOM 3713 CG ASP 394 ATOM 3714 OD1 ASP 394 ATOM 3715 OD2 ASP 394 ATOM 3716 C ASP 394 ATOM 3717 O ASP 394 ATOM 3717 O ASP 394 ATOM 3718 N ASN 395 ATOM 3720 CA ASN 395 ATOM 3721 CB ASN 395 ATOM 3721 CB ASN 395 ATOM 3722 CG ASN 395 ATOM 3722 CG ASN 395 ATOM 3723 OD1 ASN 395 ATOM 3724 ND2 ASN 395 ATOM 3727 C ASN 395 ATOM 3728 O ASN 395 ATOM 3729 N GLN 396 ATOM 3731 CA GLN 396 ATOM 3731 CA GLN 396 ATOM 3732 CB GLN 396 ATOM 3733 CG GLN 396	41.712 57.482 21.327 1.00 25.78  J8.907 58.277 23.572 1.00 J1.54  J8.885 57.086 23.916 1.00 J0.34  J8.021 58.812 22.726 1.00 J2.68  J8.021 58.812 22.726 1.00 J4.14  J7.681 56.856 21.355 1.00 J4.14  J7.681 56.352 20.184 1.00 40.62  J7.510 55.716 19.279 1.00 43.56  J7.510 55.716 19.279 1.00 43.56  J7.510 55.716 19.279 1.00 39.59  J8.440 J5.968 56.205 23.339 1.00 J4.44  J5.968 56.205 23.339 1.00 J4.44  J5.968 56.205 23.339 1.00 J4.44  J5.968 56.205 23.339 1.00 J4.65  J8.407 57.984 24.819 1.00 J6.47  J8.407 57.984 24.819 1.00 J6.47  J8.407 57.984 24.819 1.00 J6.47  J8.409 58.434 26.157 1.00 J6.56  J8.409 58.434 26.157 1.00 J6.56  J8.400 J5.14  J8.400 J5.400 J6.56  J8.400 J6.500 J6.56  J8.400 J6.500 J6.56  J8.400 J6.500 J6.500 J7.41  J8.400 J7.400 J7.41  J8.400 J7.400 J7.400  J8
ATOM 3581 CG GLU 381 ATOM 3582 CD GLU 381 ATOM 3583 OE1 GLU 381 ATOM 3584 OE2 GLU 381 ATOM 3585 C GLU 381 ATOM 3586 O GLU 381 ATOM 3587 N GLU 382 ATOM 3589 CA GLU 382 ATOM 3590 CB GLU 382 ATOM 3591 CG GLU 382 ATOM 3591 CG GLU 382 ATOM 3593 OE1 GLU 382 ATOM 3595 C GLU 382 ATOM 3595 C GLU 382 ATOM 3596 O GLU 382 ATOM 3596 O GLU 382 ATOM 3597 N GLN 383 ATOM 3599 CA GLN 383 ATOM 3600 CB GLN 383 ATOM 3600 CB GLN 383 ATOM 3601 CG GLN 383 ATOM 3603 OE1 GLN 383 ATOM 3604 NE2 GLN 383 ATOM 3606 O GLN 383 ATOM 3607 C GLN 383 ATOM 3608 O GLN 383 ATOM 3609 N LEU 384 ATOM 3611 CA LEU 384 ATOM 3612 CB LEU 384 ATOM 3613 CG LEU 384 ATOM 3614 CD1 LEU 384 ATOM 3615 CD2 LEU 384 ATOM 3616 C LEU 384 ATOM 3616 C LEU 384 ATOM 3617 O LEU 384 ATOM 3618 N GLU 385 ATOM 3618 N GLU 385 ATOM 3620 CA GLU 385	56.021 55.336 38.367 1.00 32.64 AA 55.424 55.837 39.673 1.00 31.08 AA 55.424 55.837 39.673 1.00 30.26 AA 56.134 55.337 40.915 1.00 30.26 AA 56.193 54.097 41.090 1.00 25.59 AA 56.613 56.192 41.718 1.00 28.20 AA 56.095 55.739 35.867 1.00 38.63 AA 57.201 55.342 35.467 1.00 39.43 AA 54.992 55.637 35.133 1.00 36.87 AA 54.999 55.074 33.800 1.00 34.67 AA 54.999 55.074 33.800 1.00 34.67 AA 54.998 52.888 32.572 1.00 44.29 AA 55.459 52.888 32.572 1.00 44.29 AA 55.459 51.157 33.946 1.00 51.32 AA 55.278 50.597 31.914 1.00 57.67 AA 55.278 50.597 31.914 1.00 57.67 AA 55.278 50.597 31.914 1.00 57.67 AA 54.205 56.286 31.869 1.00 37.67 AA 54.205 56.286 31.869 1.00 34.77 AA 53.274 57.090 31.071 1.00 34.77 AA 53.599 58.572 31.212 1.00 29.45 AA 54.918 60.263 32.392 1.00 24.86 AA 55.179 60.753 31.316 1.00 29.45 AA 54.918 60.263 32.392 1.00 24.86 AA 55.179 60.753 31.316 1.00 29.07 AA 55.097 60.906 33.541 1.00 28.28 AA 54.918 60.263 32.392 1.00 24.86 AA 55.179 60.753 31.316 1.00 29.07 AA 52.255 57.029 28.891 1.00 34.72 AA 54.918 60.263 32.392 1.00 24.86 AA 54.918 60.263 32.392 1.00 29.07 AA 54.391 56.216 29.126 1.00 37.41 AA 54.906 56.196 27.848 1.00 29.58 AA 54.918 56.746 27.475 1.00 32.75 AA 54.391 56.216 29.126 1.00 37.41 AA 54.906 56.196 27.848 1.00 29.58 AA 54.976 56.196 27.848 1.00 29.58 AA 54.976 56.186 27.254 1.00 28.69 AA 54.976 57.507 23.337 1.00 42.84 AA 58.755 57.507 23.337 1.00 47.19 AA 54.296 58.165 24.708 1.00 42.84 AA 54.475 57.507 23.337 1.00 47.19 AA 54.296 58.165 24	A ATOM 3739 C GLN 396 A ATOM 3740 O GLN 396 A ATOM 3741 N ASN 397 A ATOM 3743 CA ASN 397 A ATOM 3744 CB ASN 397 A ATOM 3745 CG ASN 397 A ATOM 3746 OD1 ASN 397 A ATOM 3750 C ASN 397 A ATOM 3750 C ASN 397 A ATOM 3751 O ASN 397 A ATOM 3752 N LEU 398 A ATOM 3755 CB LEU 398 A ATOM 3756 CG LEU 398 A ATOM 3756 CG LEU 398 A ATOM 3757 CD1 LEU 398 A ATOM 3758 CD2 LEU 398 A ATOM 3758 CD2 LEU 398 A ATOM 3760 O LEU 398 A ATOM 3761 N GLN 399 A ATOM 3761 N GLN 399 A ATOM 3766 CD GLN 399 A ATOM 3767 OE1 GLN 399 A ATOM 3766 CD GLN 399 A ATOM 3770 CE GLN 399 A ATOM 3771 C GLN 399 A ATOM 3771 C GLN 399 A ATOM 3776 CB GLN 400 A ATOM 3777 CG GLN 400 A ATOM 3778 CD GLN 400 A ATOM 3778 CD GLN 400 A ATOM 3779 OE1 GLN 400 A ATOM 3780 NE2 GLN 400 A ATOM 3780 NE2 GLN 400	30.019 55.422 19.535 1.00 42.50 AAAA 29.959 59.029 24.292 1.00 38.09 AAAA 29.418 60.055 23.913 1.00 39.65 AAAA 28.794 59.115 26.467 1.00 38.64 AAAA 27.795 58.035 26.846 1.00 35.69 AAAA 27.795 58.035 26.846 1.00 39.95 AAAA 27.182 57.370 25.624 1.00 39.95 AAAA 27.484 56.084 25.417 1.00 36.45 AAAA 29.264 59.795 27.750 1.00 38.08 AAAA 29.264 59.795 27.750 1.00 38.08 AAAA 29.264 59.795 27.750 1.00 38.08 AAAA 23.725 60.452 29.099 1.00 38.24 AAAA 33.536 59.791 30.199 1.00 38.47 AAAA 33.536 59.791 30.199 1.00 33.50 AAAA 34.095 58.432 29.882 1.00 35.71 AAAA 34.095 58.432 29.882 1.00 35.71 AAAA 30.879 61.912 29.361 1.00 40.51 AAAA 30.879 61.912 29.361 1.00 44.82 AAAA 29.743 63.498 30.458 1.00 44.82 AAAA 29.743 63.498 30.845 1.00 46.98 AAAA 29.743 63.498 30.458 1.00 63.27 AAAA 29.744 63.515 32.919 1.00 42.85 AAAA 29.744 63.515 32.919 1.00 42.85 AAAA 20.410 62.2188 38.095 1.00 43.73 AAAA 20.440 62.2188 38.095 1.00 45.70 AAAA 20.440 62.2188 38.095 1.00 44.99 AAAA 20.440 62.2188 38.095 1.00 44.99 AAAA 20.440 62.2188 38.095 1.00 44.99 AAAA 20.440
ATOM 3622 CG GLU 385 ATOM 3623 CD GLU 385 ATOM 3624 OE1 GLU 385 ATOM 3625 OE2 GLU 385 ATOM 3625 OE2 GLU 385 ATOM 3626 C GLU 385 ATOM 3626 C GLU 385 ATOM 3626 C GLU 385 ATOM 3627 O GLU 385 ATOM 3628 N GLY 386 ATOM 3631 C GLY 386 ATOM 3631 C GLY 386 ATOM 3631 C GLY 386 ATOM 3632 O GLY 386 ATOM 3635 CA ASN 387 ATOM 3636 CB ASN 387 ATOM 3638 OD1 ASN 387 ATOM 3638 OD1 ASN 387 ATOM 3639 ND2 ASN 387 ATOM 3644 C ASN 387 ATOM 3644 N TYR 388 ATOM 3646 CA TYR 388 ATOM 3646 CA TYR 388 ATOM 3650 CE1 TYR 388 ATOM 3650 CE1 TYR 388 ATOM 3651 CC TYR 388 ATOM 3652 CE2 TYR 388 ATOM 3656 C TYR 388 ATOM 3657 O TYR 388 ATOM 3666 CA SER 389 ATOM 3667 C TYR 388 ATOM 3667 C TYR 388 ATOM 3668 CA PHE 390 ATOM 3667 C PHE 390 ATOM 3667 C PHE 390 ATOM 3667 C PHE 390 ATOM 3670 CC PHE 390 ATOM 3671 CD1 PHE 390 ATOM 3675 CZ PHE 390 ATOM 3676 C PHE 390 ATOM 3677 O PHE 390 ATOM 3678 C PHE 390 ATOM 3678 C PHE 390 ATOM 3679 CC PHE 390 ATOM 3679 CC PHE 390 ATOM 3678 N TYR 391 ATOM 3688 C CD TYR 391 ATOM 3689 C TYR 391 ATOM 3699 C TYR 391	\$5.273 \$6.189 23.393 1.00 \$3.13 AAA \$4.464 \$4.951 23.875 1.00 \$8.73 AAA \$53.470 \$55.066 24.640 1.00 \$7.99 AAA \$53.680 \$9.551 24.625 1.00 42.49  \$52.469 \$59.694 24.402 1.00 44.93 AAA \$4.621 \$60.561 24.827 1.00 39.27 \$4.062 \$61.936 24.819 1.00 36.93 AAA \$54.062 \$61.936 24.819 1.00 36.93 AAA \$54.012 \$63.490 26.665 1.00 34.95 \$4.458 \$61.299 27.066 1.00 36.01 AAA \$54.052 \$61.299 27.066 1.00 36.01 AAA \$54.052 \$61.299 27.066 1.00 36.01 AAA \$55.608 \$62.377 28.828 1.00 39.59 AAA \$55.608 \$62.377 28.828 1.00 39.59 AAA \$55.980 \$61.681 28.685 1.00 44.06 AAA \$58.057 \$62.475 28.650 1.00 44.07 AAA \$53.123 \$61.880 29.055 1.00 37.24 AAA \$53.123 \$61.880 29.055 1.00 37.24 AAA \$53.123 \$61.880 29.055 1.00 37.24 AAA \$52.972 \$63.003 29.522 1.00 38.63 AAA \$9.921 \$62.177 27.300 1.00 27.49 AAA \$49.864 \$60.982 28.267 1.00 27.49 AAA \$49.921 \$62.177 27.300 1.00 27.26 AAA \$49.931 \$63.417 27.707 1.00 28.46 AAA \$49.645 \$64.499 26.873 1.00 24.65 AAA \$49.645 \$64.499 26.873 1.00 24.65 AAA \$49.645 \$64.499 26.873 1.00 24.65 AAA \$49.645 \$64.499 26.873 1.00 24.98 AAA \$49.645 \$64.499 26.873 1.00 25.98 AAA \$50.124 \$65.403 24.786 1.00 33.18 AAA \$49.645 \$64.499 26.873 1.00 24.65 AAA \$49.645 \$64.499 26.873 1.00 24.95 AAA \$49.645 \$64.499 26.873 1.00 34.00 AAA \$49.645 \$64.499 26.873 1.00 25.98 AAA \$49.645 \$64.499 26.873 1.00 25.98 AAA \$49.645 \$64.499 26.873 1.00 26.92 AAA \$49.645 \$64.499 26.873 1.00 25.98 AAA \$49.645 \$64.499 26.873 1.00 34.00 AAA \$49.645 \$64.499 26.873 1.00 34.00 AAA \$49.645 \$64.499 26.873 1.00 34.00 AAA \$49.645 \$64.600 560 34.00 AAA \$40.645 \$69.600 34.200 30.000 34.30 AAA \$40.646 \$60.950 32.890 30.000 30.300 AAA \$40.647 \$69.950 32.890 30.000 30.300 AAA \$40.640 \$60.950 32.890 30.000 30.300 30.300 AA	A ATOM 3784 O GLN 400 A ATOM 3785 N LEU 401 A ATOM 3787 CA LEU 401 A ATOM 3788 CB LEU 401 A ATOM 3788 CB LEU 401 A ATOM 3789 CG LEU 401 A ATOM 3799 CG LEU 401 A ATOM 3791 CD2 LEU 401 A ATOM 3791 CD2 LEU 401 A ATOM 3793 O LEU 401 A ATOM 3794 N TRP 402 A ATOM 3796 CA TRP 402 A ATOM 3796 CA TRP 402 A ATOM 3798 CG TRP 402 A ATOM 3799 CD2 TRP 402 A ATOM 3799 CD2 TRP 402 A ATOM 3799 CD2 TRP 402 A ATOM 3800 CE2 TRP 402 A ATOM 3801 CE3 TRP 402 A ATOM 3801 CE3 TRP 402 A ATOM 3802 CD1 TRP 402 A ATOM 3805 CZ2 TRP 402 A ATOM 3806 C TRP 402 A ATOM 3807 CH2 TRP 402 A ATOM 3808 C TRP 402 A ATOM 3808 C TRP 402 A ATOM 3810 N ASP 403 A ATOM 3811 CB ASP 403 A ATOM 3813 CB ASP 403 A ATOM 3813 CB ASP 403 A ATOM 3816 OD2 ASP 403 A ATOM 3816 OD2 ASP 403 A ATOM 3817 C ASP 403 A ATOM 3816 CD2 ASP 403 A ATOM 3817 C ASP 403 A ATOM 3818 O ASP 403 A ATOM 3819 N TRP 404 A ATOM 3821 CA TRP 404 A ATOM 3822 CB TRP 404 A ATOM 3823 C TRP 404 A ATOM 3824 O TRP 404 A ATOM 3829 CA ASP 405 A ATOM 3830 CA ARG 407 A ATOM 3831 CA ARG 407 A ATOM 3831 CA ARG 407 A ATOM 3836 CA ARG 407 A A	32.992 61.953 34.111 1.00 45.39 AAAA 34.059 61.751 34.823 1.00 41.97 AAAA 35.177 62.941 35.243 1.00 41.07 AAAA 36.445 63.813 35.231 1.00 38.05 AAAA 36.800 64.425 33.857 1.00 31.72 AAAA 37.123 63.330 32.887 1.00 27.88 AAAA 34.806 62.415 36.639 1.00 40.74 AAAA 34.224 61.335 36.773 1.00 39.04 AAAA 35.128 63.203 37.659 1.00 40.74 AAAA 35.128 63.203 37.659 1.00 43.32 AAAA 36.806 62.456 39.891 1.00 48.15 AAAA 36.806 62.458 39.891 1.00 48.15 AAAA 36.784 61.226 39.735 1.00 39.94 37.854 60.88 38.833 1.00 44.37 AAAA 38.201 59.545 39.072 1.00 45.95 AAAA 38.553 61.609 37.846 1.00 39.31 AAAA 38.201 59.545 39.072 1.00 44.07 AAAA 39.220 58.901 88.354 1.00 44.07 AAAA 39.220 58.901 80.354 1.00 51.68 39.543 60.977 37.144 1.00 38.52 AAAA 39.674 64.271 39.990 1.00 65.12 AAAA 31.216 64.771 39.994 1.00 51.62 AAAA 31.226 64.951 40.283 1.00 58.06 AAAA 31.236 64.714 39.990 1.00 65.12 AAAA 31.266 64.74 49.930 1.00 66.24 AAAA 31.266 64.951 40.283 1.00 58.06 AAAA 31.276 66.954 41.31 1.00 59.06 AAAA 31.281 67.555 42.010 1.00 67.88 AAAA 31.296 66.977 40.805 1.00 61.21 AAAA 31.297 66.379 41.31 1.00 77.66 AAAA 31.297 66.379 41.31 1.00 78.46 AAAA 31.299 66.297 40.805 1.00 61.21 AAAA 31.296 66.375 43.038 1.00 77.29 AAAA 31.297 66.872 41.314 1.00 78.16 AAAA 31.299 66.827 41.314 1.00 79.53 AAAA 31.299 66.375 43.038 1.00 77.29 AAAA 31.299 66.817 44.4198 1.00 79.53 AAAA 31.299 66.817 44.688 1.00 77.29 AAAA 31.299 66.817 44.688 1.00 77.29 AAAA 31.399 65.417 69.899 1.00 66.78 AAAA 31.399 65.417 69.899 1.00 66.78 AAAA 31.399 65.541 41.316 1.00 60.54 AAAA 31.299 66.817 44.688 1.00 77.29 AAAA 31.990 66.817 44.688 1.00 77.29 AAAA 31.990 66.817 44.688 1.00 77.29 AAAA 31.990 66.817 44.689 1.00 77.29 AAAA 31.990 66.817 44.689 1.00 77.29 AAAA 31.990 66.817 44.650 1.00 67.78 AAAA 31.990 66.817 44.650 1.00 67.78 AAAA 31.990 66.817 44.650 1.00 77.25 AAAA 31.990 66.817 44.650 1.00 77.25 AAAA 31.990 66.817 44.650 1.00

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•	ATOM MOTA		409 44 410 45	.317 63.893	42.316 42.198	1.00 48.67 1.00 48.82	AAAA AAAA AAAA	MOTA MOTA MOTA	4031 O VAL 4032 N SER 4034 CA SER	426 427 427	34.946 34.695	72.579 73.194	30.000 31.293	1.00 45.01 1.00 46.62 1.00 48.46	**** **** ****
	A HOTA ATOM ATOM	3876 CA THR 3877 CB THR 3878 OG1 THR 3880 CG2 THR 3881 C THR	410 47 410 46 410 48	.364 64.730 .860 65.897 .861 64.881 .506 63.740	43.270 43.940 42.997 41.009	1.00 46.46 1.00 47.45 1.00 48.99 1.00 47.61 1.00 43.42	AAAA AAAA AAAA AAAA	MOTA MOTA MOTA MOTA MOTA	4035 CB SER 4036 OG SER 4038 C SER 4039 O SER 4040 N GLU	427 427 427 427 428	33.184 32.597 35.237 35.928 34.946	73.447 73.899 72.316 72.794	31.456 30.229 32.421 33.318	1.00 49.20 1.00 51.43 1.00 48.91 1.00 49.65 1.00 49.00	AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM ATOM	3882 O THR 3883 N ILE 3885 CA ILE 3886 CB ILE 3887 CG2 ILE	-411 48 411 48 411 48	.035 64.406 .934 63.735	39.995 39.070 37.569	1.00 42.61 1.00 41.25 1.00 41.84 1.00 36.01 1.00 31.64	AAAA AAAA AAAA AAAA	MOTA MOTA MOTA MOTA	4042 CA GLU 4043 CB GLU 4044 CG GLU 4045 CD GLU 4046 OE1 GLU	428 428 428 428 428	35.425 34.693 33.197 32.829 32.667	70.123 68.787 68.875 69.284	33.402 33.309 33.541 34.969	1.00 50.84 1.00 53.24 1.00 60.59 1.00 65.08	**** **** ****
	ATOM ATOM ATOM ATOM	3888 CG1 ILE 3889 CD1 ILE 3890 C ILE 3891 O ILE	411 47 411 46 411 50	.050 63.889 .594 62.884 .338 64.265 .730 65.375	37.375 36.336 39.407	1.00 36,00 1.00 31.18 1.00 45.51	AAAA AAAA AAAA	ATOM ATOM ATOM	4047 OE2 GLU 4048 C GLU 4049 O GLU	428 428 428 429	32.696 36.921 37.706 37.315	70.514 69.886 69.999	35.209 33.234 34.168	1.00 66.54 1.00 68.29 1.00 51.94 1.00 56.08	***** ***** *****
	ATOM ATOM ATOM	3892 N LYS 3894 CA LYS 3895 CB LYS 3896 CG LYS 3898 C LYS	412 52 412 52 412 53	.419 63.769 .972 62.563	40.634 41.369 42.746	1.00 43.71 1.00 41.26 1.00 42.01 1.00 49.93 1.00 41.15	AAAA AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4052 CA ILE 4053 CB ILE 4054 CG2 ILE 4055 CG1 ILE 4056 CD1 ILE	429 429 429 429 429	38.697 38.897 40.359 38.214	69.136 69.094 67.854	30.240 29.893 29.785	1.00 48.09 1.00 45.91 1.00 43.32 1.00 42.83 1.00 44.47	AAAA AAAA AAAA
	ATOM ATOM ATOM ATOM	3899 O LYS 3900 N ALA 3902 CA ALA 3903 CB ALA	412 54 413 53 413 54	.195 64.959 .192 63.308 .043 63.445 .277 62.614	39.560 38.411 37.255 37.471	1.00 40.96 1.00 40.81 1.00 40.95 1.00 44.44	AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4057 C ILE 4058 O ILE 4059 N TYR 4061 CA TYR	429 429 430 430	38.390 39.645 40.722 39.268 40.189	70.319 69.965 71.596	32,332 32.817 32.312	1.00 50.57 1.00 50.48 1.00 53.59 1.00 56.94	AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM	3904 C ALA 3905 O ALA 3906 N GLY 3908 CA GLY	413 52 414 53 414 53	.599 62.071 .694 63.729 .086 63.434	35.929 34.884 33.603	1.00 40.18 1.00 40.51 1.00 39.10 1.00 40.43 1.00 41.42	AAAA AAAA AAAA	ATOM ATOM ATOM	4062 CB TYR 4063 CG TYR 4064 CD1 TYR 4065 CE1 TYR	430 430 430 430	40.000 39.958 39.656 39.447	74.055 72.923 72.997	30.692 29.943 28.583	1.00 59.37 1.00 61.64 1.00 61.92 1.00 61.98	AAAA AAAA AAAA
	ATOM ATOM ATOM ATOM	3909 C GLY 3910 O GLY 3911 N LYS 3913 CA LYS 3914 CB LYS	414 51 415 51 415 50	.870 65.415 .161 64.247 .065 65.173	34.131 32.347 32.040	1.00 42.72 1.00 40.85 1.00 38.88 1.00 39.32	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4066 CD2 TYR 4067 CE2 TYR 4068 CZ TYR 4069 OH TYR 4071 C TYR	430 430 430 430 430	40.073 39.863 39.539 39.226 40.068	74.206 74.254	28.641 27.935 26.595	1.00 62.01 1.00 61.34 1.00 61.30 1.00 62.51 1.00 55.10	AAAA AAAA AAAA AAAA
	MOTA MOTA MOTA MOTA	3915 CG LYS 3916 CD LYS 3917 CE LYS 3918 NZ LYS 3922 C LYS	415 51 415 52 415 52	.618 66.153 .669 66.925	28.746 28.048 26.813	1.00 40.34 1.00 38.47 1.00 35.12 1.00 43.52 1.00 37.63	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4072 O TYR 4073 N ARG 4075 CA ARG 4076 CB ARG 4077 CG ARG	430 431 431 431 431	40.961 38.964 38.776 37.377 37.349	72.229 72.253 71.782	34.908 36.346 36.742	1.00 56.49 1.00 52.81 1.00 52.51 1.00 54.84 1.00 57.04	AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM	3923 O LYS 3924 N MET 3926 CA MET 3927 CB MET	415 48 416 47 416 46 416 45	.834 63.201 .962 65.234 .759 64.690 .548 65.090	31.334 30.834 30.204 31.032	1.00 37.93 1.00 35.60 1.00 30.58 1.00 31.10	AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4079 C ARG 4080 O ARG 4081 N MET 4083 CA MET	431 431 432 432	39.770 40.703 39.536 40.376	71.250 71.582 70.009 68.866	36.838 37.554 36.427 36.766	1.00 51.03 1.00 49.10 1.00 52.81 1.00 55.38	AAAA AAAA AAAA
	ATOM ATOM ATOM	3928 CG MET 3929 SD MET 3930 CE MET 3931 C MET 3932 O MET	416 44 416 44 416 46	.474 65.039	33.413 33.433 28.755		AAAA AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM ATOM	4084 CB MET 4085 CG MET 4085 SD MET 4087 CE MET 4088 C MET	432 432 432 432 432		66.348 65.457 63.999	35.402 35.365 34.999	1.00 52.88 1.00 50.22 1.00 47.27 1.00 42.56 1.00 58.02	**************************************
	ATOM ATOM ATOM ATOM	3933 N TYR 3935 CA TYR 3936 CB TYR 3937 CG TYR	417 45 417 45 417 46 417 45	.797 64.114 .375 64.284 .137 63.349 .660 63.489	28.090 26.710 25.779 24.350	1.00 24.40 1.00 25.54 1.00 25.59 1.00 29.15	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4089 O MET 4090 N GLU 4092 CA GLU 4093 CB GLU	432 433 433 433	42.710 42.131 43.480 43.447	68.694 70.161 70.643 71.636	37.346 35.732 35.483 34.349	1.00 59.47 1.00 59.97 1.00 60.19 1.00 60.90	4444 4444 4444 4444
	MOTA MOTA MOTA MOTA MOTA	3938 CD1 TYR 3939 CE1 TYR 3940 CD2 TYR 3941 CE2 TYR 3942 CZ TYR	417 45 417 44 417 44	.383 64.867 .967 62.453 .479 62.622	22.368 23.697 22.382	1.00 27.78 1.00 28.95 1.00 29.30 1.00 26.16 1.00 28.87	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM ATOM	4094 CG GLU 4095 CD GLU 4096 CE1 GLU 4097 CE2 GLU 4098 C GLU	433 433 433 433 433	43.854 44.322 44.683 44.325 44.032	72.242 73.282 72.117	32.159 32.742 30.920	1.00 62.57 1.00 64.14 1.00 64.05 1.00 66.58 1.00 60.66	**************************************
	MOTA ! MOTA ! MOTA ! MOTA !	3943 OH TYR 3945 C TYR 3946 O TYR 3947 N PHE 3949 CA PHE	417 43. 417 43. 418 43.	.877 63.956 .443 62.845 .081 64.917	26.615 26.953 26.173	1.00 38.55 1.00 29.17 1.00 30.30 1.00 30.27 1.00 33.08	8848 8848 8848	ATOM ATOM ATOM ATOM ATOM	4099 O GLU 4100 N GLU 4102 CA GLU 4103 CB GLU 4104 CG GLU	433 434 434 434 434	43.147 43.541 42.502	72.035 72.770 73.838	37.416 38.603 38.923	1.00 62.11 1.00 60.35 1.00 60.47 1.00 63.91 1.00 67.95	AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM	3950 CB PHE 3951 CG PHE 3952 CD1 PHE	418 40. 418 40.	.869 65.640	26.958 28.393	1.00 32.78 1.00 32.46	AAAA AAAA	ATOM ATOM ATOM	4105 CD GLU 4106 OE1 GLU 4107 OE2 GLU	434 434 434	40.430 39.629	75.127 74.759	38.224 39.082	1.00 72.47 1.00 75.10 1.00 75.28	AAAA AAAA AAAA
	MCTA MOTA MOTA MOTA	3953 CD2 PHE 3954 CE1 PHE 3955 CE2 PHE 3956 CZ PHE 3957 C PHE	418 41. 418 39. 418 40.	.771 64.024	30.636 30.204 31.090	1.00 31.60 1.00 34.33 1.00 33.90	4444 4444 4444 4444 4444	ATCM ATOM ATOM ATCM ATOM	4103 C GLU 4109 O GLU 4110 N VAL 4112 CA VAL 4111 CB VAL	434 434 435 435 435		72.185 70.834 69.943	40.667 39.936 41.081	1.00 58.76 1.00 59.92 1.00 55.65 1.00 53.00 2.00 51.91	4444 4444 4444 4444 4444
	MOTA : MOTA : MOTA : MOTA :	3958 O PHE 3959 N ALA 3961 CA ALA 3962 CB ALA	418 41. 419 40. 419 39. 419 41.	.532 65.955 .439 63.976 .970 64.124 .140 64.071	24.019 24.047 22.675 21.736	1.00 35.67 1.00 33.64 1.00 35.34 1.00 32.63	AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4114 CG1 VAL 4115 CG2 VAL 4116 C VAL 4117 O VAL	435 435 435 435	41.674 40.525 44.194 44.944	68.524 70.052 68.954 68.838	42.647 41.031 41.018 41.988	1.00 50.00 1.00 50.48 1.00 52.57 1.00 55.93	4444 4444 4444 4444
	ATOM ATOM : ATOM : ATOM : ATOM	3963 C ALA 3964 O ALA 3965 N PHE 3967 CA PHE 3968 CB PHE	419 39. 420 37. 420 36.	.136 61.888 .827 63.527 .781 62.633	22.418 21.675 21.177	1.00 38.10 1.00 42.66 1.00 35.56 1.00 35.67 1.00 35.36	AAAA AAAA AAAA AAAA	ATOM MOTA ATOM ATOM	4118 N THR 4120 CA THR 4121 CB THR 4122 OG1 THR 4124 CG2 THR	436 436 436 436 436	44.339 45.400 45.309 44.904 44.295	67.285 66.575 67.511	39.767 38.432 37.436	1.00 49.37 1.00 46.23 1.00 43.66 1.00 41.65 1.00 44.73	AAAA AAAA AAAA AAAA
(	MOTA M	3969 CG PHE 3970 CD1 PHE 3971 CD2 PHE 3972 CE1 PHE 3973 CE2 PHE	420 39. 420 36. 420 39.	.817 61.756 .165 61.782 .911 62.227	19.039 18.732 18.100 17.517	1.00 35.13 1.00 33.79 1.00 35.56 1.00 33.63	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM	4125 C THR 4126 O THR 4127 N GLY 4129 CA GLY	436 436 437 437	46.750 47.751 46.760 47.980	67.368 69.254 70.036	40.282 39.527 39.572	1.00 47.54 1.00 47.63 1.00 48.49 1.00 48.06 1.00 48.65	AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM	3974 CZ PHE 3975 C PHE 3976 O PHE 3977 N ASN	420 38. 420 35. 420 35.	706 62.771	16.608 22.181 22.133	1.00 33.07 1.00 38.38 1.00 40.78	AAAA AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4130 C GLY 4131 O GLY 4132 N THR 4134 CA THR 4135 CB THR	437 438 438 438	48.713 49.854 48.053 48.623 47.899	70.392 69.311 69.129 68.018	38.122 37.265 35.933 35.180	1.00 48.33 1.00 46.21 1.00 45.29 1.00 45.26	AAAA AAAA AAAA AAAA
	MOTA MOTA MOTA MOTA MOTA	3979 CA ASN 3980 CB ASN 3981 CG ASN 3982 OD1 ASN 3983 ND2 ASN	421 34. 421 36. 421 36.	.400 62.821 .952 63.077 .248 62.374 .294 61.142 .323 63.147	25.527 25.770 25.946	1.00 33.41 1.00 29.98 1.00 25.86	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4136 OG1 THR 4138 CG2 THR 4139 C THR 4140 O THR 4141 N LYS	438 438 438 438 439	48.953 48.529 49.050	66.857 70.425 70.553	34.895 35.148 34.039	1.00 47.25 1.00 45.44 1.00 43.52 1.00 42.01 1.00 44.19	**************************************
	ATOM ATOM ATOM	3986 C ASN 3987 O ASN 3988 N PRO 3989 CD PRO	421 33. 421 33. 422 32. 422 32.	.379 63.901 .456 65.031 .430 63.581 .213 62.321	23.775 24.257 22.885 22.160	1.00 39.11 1.00 36.50 1.00 39.75 1.00 36.04	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM	4143 CA LYS 4144 CB LYS 4145 CG LYS 4147 C LYS	439 439 439 439	47.722 47.171 46.711 49.114	72.683 73.701 74.873 73.096	35.115 36.114 35.455 34.689	1.00 47.05 1.00 46.93 1.00 46.45 1.00 48.32	**************************************
	ATOM ATOM ATOM ATOM ATOM	3990 CA PRO 3991 CB PRO 3992 CG PRO 3993 C PRO 3994 O PRO	422 30. 422 30. 422 30.	.451 64.615 .423 63.873 .776 62.403 .805 65.306 .664 66.536	21.680 21.781 23.716	1.00 37.57 1.00 37.38 1.00 41.51	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4148 O LYS 4149 N GLY 4151 CA GLY 4152 C GLY 4153 O GLY	439 440 440 440	49.285 50.601 51.444 52.146	73.493 73.941 72.925 73.254	33.439 33.024 32.294 31.347	1.00 51.40 1.00 49.69 1.00 49.95 1.00 50.27 1.00 50.04	8888 8888 8888 8888
	MOTA MOTA MOTA MOTA	3995 N LYS 3997 CA LYS 3998 CB LYS 3999 CG LYS 4000 CD LYS	423 29. 423 28. 423 27.	.456 64.507 .751 65.012 .682 64.002 .802 63.532 .340 62.094	25.888 26.303 25.157	1.00 45.63 1.00 47.46 1.00 53.04	AAAA AAAA AAAA AAAA	MOTA MOTA MOTA MOTA	4154 N ARG 4156 CA ARG 4157 CB ARG 4158 CG ARG 4159 CD ARG	441 441 441 441	52.195 52.550 51.411	70.712 69.574 69.098	32.021 32.956 33.777	1.00 51.02 1.00 52.95 1.00 51.15 1.00 48.21 1.00 49.62	**************************************
	MOTA MOTA MOTA MOTA	4001 CE LYS 4002 NZ LYS 4006 C LYS 4007 O LYS	423 25. 423 25. 423 30. 423 29.	.846 62.011 .448 60.689 .560 65.392 .991 65.629	25.745 26.353 27.108 28.165	1.00 63.87 1.00 64.25 1.00 45.01 1.00 45.99	AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4160 NE ARG 4152 CZ ARG 4163 NH1 ARG 4166 NH2 ARG	441 441 441	52.149 52.437 52.508 52.649	68.366 67.530 66.220 68.019	36.002 36.987 36.757 18.197	1.00 46.95 1.00 42.94 1.00 40.41 1.00 45.43	AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM ATOM ATOM	4008 N LEU 4010 CA LEU 4011 CB LEU 4012 CG LEU 4013 CD1 LEU	424 32. 424 34. 424 35.	.875 65.463 .706 65.813 .176 65.630 .110 65.328 .334 66.579	28.108 27.772 28.938	1.00 44.01 1.00 43.31 1.00 45.34	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4169 C ARG 4170 O ARG 4171 N GLN 4173 CA GLN 4174 CB GLN	441 441 442 442 442	51.341 51.074 50.380	58.964 71.048 70.600	30.678 29.891 28.673	1.00 55.03 1.00 57.67 1.00 58.46 1.00 62.79 1.00 61.21	AAAA AAAA AAAA
	MOTA MOTA MOTA MOTA	4014 CD2 LEU 4015 C LEU 4016 O LEU 4017 N CYS 4019 CA CYS	424 34. 424 32. 424 31. 425 33.	495 64.240 445 67.245 647 67.495 109 68.186 939 69.595	29.780 28.512 29.413 27.860	1.00 47.64 1.00 44.69 1.00 49.30 1.00 42.99	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM ATOM	4175 CG GLN 4176 CD GLN 4177 OE1 GLN 4178 NE2 GLN 4181 C GLN	442 442 442 442 442	48.349 47.687 47.919 46.850	70.991 69.771 68.645 69.988	30.196 30.787 30.344 31.792	1.00 62.46 1.00 60.41 1.00 59.88 1.00 60.57 1.00 65.82	AAAA AAAA AAAA AAAA
	ATOM ATOM ATOM ATOM	4020 C CYS 4021 O CYS 4022 CB CYS 4023 SG CYS	425 34. 425 34. 425 31. 425 31.	.251 70.164 .611 69.927 .843 69.752 .239 71.440	28.727 29.867 29.334 29.429	1.00 42.97 1.00 41.72 1.00 49.81 1.00 70.31	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4182 O GLN 4183 N SER 4185 CA SER 4186 CB SER	442 443 443 443	51.588 50.228 50.521 51.164	72.294 70.943 71.562 70.532	27.478 26.295 25.014 24.076	1.00 69.50 1.00 67.54 1.00 67.01 1.00 67.85	2222 2222 2222 2222
	ATOM ATOM ATOM ATOM ATOM	4024 N VAL 4026 CA VAL 4027 CB VAL 4028 CG1 VAL 4029 CG2 VAL	426 36. 426 36. 426 37. 426 35.	.963 70.924 .243 71.486 .716 72.578 .128 71.951 .619 73.595	28.308 27.327 25.999 27.132	1.00 44.40 1.00 44.04 1.00 40.82 1.00 46.91	AAAA AAAA AAAA AAAA	ATOM ATOM ATOM ATOM	4187 C SER 4188 O SER 4189 N LYS 4191 CA LYS 4192 CB LYS	443 444 444 444	48.364 49.132 47.974 48.439	72.622 72.010 72.523 71.105	25.107 23.081 22.365 21.027	1.00 67.28 1.00 67.90 1.00 67.13 1.00 66.69 1.00 67.24	AAAA AAAA AAAA AAAA
	ATOM	4030 C VAL	426 )6.	131 72.083	₹9.82	1.00 44.63	AAAA	ATOM	4193 C LYS	444	40.808	11.354	45/	1.00 65.75	****

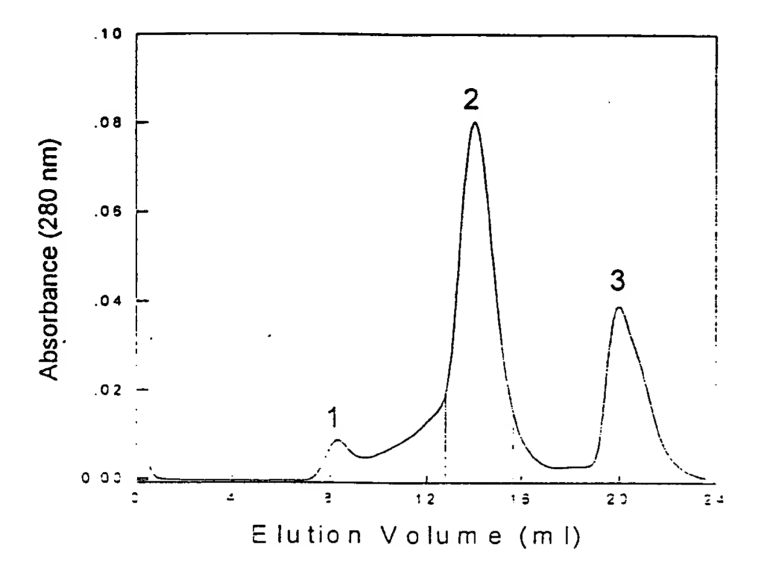
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ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4194 O LYS 4195 N GLY 4197 CA GLY 4198 C GLY 4199 O GLY 4200 N ASP 4202 CA ASP 4203 CB ASP 4204 CG ASP 4205 OD1 ASP 4206 OD2 ASP 4207 C ASP 4208 O ASP 4209 N ILE 4211 CA ILE 4212 CB ILE 4211 CA ILE 4212 CB ILE 4213 CG2 ILE 4214 CG1 ILE 4215 CD1 ILE 4216 C ILE 4217 O ILE 4218 N ASN 4220 CA ASN 4221 CB ASN 4221 CB ASN 4222 CG ASN 4223 OD1 ASN 4224 ND2 ASN 4227 C ASN 4228 O ASN 4227 C ASN 4231 CA THR 4232 CB THR 4233 CG1 THR 4233 CG1 THR 4235 CG2 THR 4236 C THR 4237 O THR 4238 N ARG 4240 CA ARG 4241 CB ARG 4237 O THR 4238 N ARG 4240 CA ARG 4241 CB ARG 4241 CB ARG 4242 CG ARG 4241 CB ARG 4242 CG ARG 4255 N ASN 4256 C ARG 4257 CA ASN 4258 CB ASN 4259 CG ASN 4260 OD1 ASN 4261 ND2 ASN 4257 CA ASN 4258 CB ASN 4259 CG ASN 4260 OD1 ASN 4261 ND2 ASN 4262 C ARG 4270 C ASN 4263 C ARG 4271 O ASN 4264 C ASN 4265 O ASN 4266 N ASN 4267 CB ASN 4268 CA ASN 4269 CB ASN 4269 CB ASN 4269 CB ASN 4270 C ASN 4271 O ASN 4271 O ASN 4272 N GLY 4275 C GLY 4276 O GLY 4277 N GLU 4279 CA GLU	447 43.069 447 41.916 447 41.633 448 41.662 448 41.019 448 41.106 448 40.575 448 39.408 448 39.572 449 39.134 449 37.781 449 37.815 449 38.913 449 36.981 449 37.785 450 37.658 450 37.658 450 37.658 450 41.143 450 450 41.612 450 41.143 450 450 450 450 450 450 450 450 450 450	70.421 21.445 69.423 21.117 68.522 22.247 67.610 22.064 68.806 23.414 68.099 24.668 68.767 25.819 68.141 26.059 67.456 25.174 68.350 27.162 68.197 24.955 67.202 25.011 69.423 25.115 69.646 25.407 70.389 26.758 70.386 27.225 70.172 29.231 70.438 24.292 71.632 24.450 69.790 23.156 70.516 22.056 69.766 20.718 68.378 20.800 67.421 20.347 68.246 21.395 70.820 22.370 70.044 23.027 71.970 21.892 72.433 22.423 74.629 21.212 74.198 23.479 72.185 20.880 72.445 20.812 71.655 19.885 71.401 18.632 71.755 17.483 70.561 16.801 70.955 16.073 70.821 16.942	1.00 63.29 1.00 59.11 1.00 55.66 1.00 52.13 1.00 53.96 1.00 49.82 1.00 46.80 1.00 43.77 1.00 39.45 1.00 45.58 1.00 45.58 1.00 44.37 1.00 43.79 1.00 44.45 1.00 43.79 1.00 41.47 1.00 41.47 1.00 41.47 1.00 55.78 1.00 55.78 1.00 55.78 1.00 55.89 1.00 55.89 1.00 55.89 1.00 55.89 1.00 55.89 1.00 55.89 1.00 57.21 1.00 59.40 1.00 55.58 1.00 57.21 1.00 59.40 1.00 52.99 1.00 52.99 1.00 52.99 1.00 53.18 1.00 52.99 1.00 53.18 1.00 52.79 1.00 46.87 1.00 47.57 1.00 45.63 1.00 46.87 1.00 46.87 1.00 46.87 1.00 47.57 1.00 48.17 1.00 46.87 1.00 57.18 1.00 60.27 1.00 59.45 1.00 64.99 1.00 57.87 1.00 64.99 1.00 64.99 1.00 64.99 1.00 64.99 1.00 64.99 1.00 64.99 1.00 64.99 1.00 64.99 1.00 64.99 1.00 64.98 1.00 64.99 1.00 67.97 1.00 69.95 1.00 70.80 1.00 70.80 1.00 70.80 1.00 74.27 1.00 74.08 1.00 76.62	**************************************	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4392 O3 k 4394 C4 k 4396 O4 k 4398 C5 k 4400 O5 k 4401 C6 k 4408 C2 k 4410 N2 k 4410 C7 k 4413 C8 k 4412 C7 k 4413 C8 k 4422 C4 k 4427 O5 k 4428 C6 k 4431 O6 k 4432 C2 k 4437 N2 k 4439 C7 k 4439 C7 k 4439 C7 k 4449 C4 k 4451 O4 k 4452 C5 k 4454 O5 k 4452 C5 k 4454 O5 k 4455 C6 k 4460 C1 k 4452 C5 k 4454 O5 k 4457 O5 k 4460 C1 k 4452 C5 k 4453 C2 k 4453 C3 k 4454 C5 k 4458 C6 k 4469 C4 k 4460 C1 k 4460 C2 k 4461 C3 k 4461 C3 k 4462 C2 k 4463 C3 k 4463 C3 k 4464 C3 k 4465 C5 k 4469 C4 k 4469 C4 k 4469 C4 k 4469 C4 k 4469 C5 k 4469 C6 k 4469 C	IAG 105B IAG 284A IAG 284B IAG	29.721 29.6330 30.4154 30.325 56.325 56.893 56.895 57.275 56.895 57.275 56.895 57.806	21.268 21.998 19.899 19.899 19.950 45.501 46.6689 48.1720 45.1669 48.1320 45.1720 45.45.1720 45.45.1720 45.45.1720 45.45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 45.1720 47.1720	75.373 76.799 77.78.007 78.007 78.007 78.007 78.007 78.007 78.007 78.007 78.007 66.584 67.331 69.912 69.913 67.593 69.913 67.686 67.866 67.866 67.866 77.77 77.866 77.77 77.666 77.77 77.666 77.77	1.00 65.10 1.00 72.19 1.00 65.51 1.00 65.49 1.00 64.84 1.00 62.03 1.00 67.68 1.00 43.05 1.00 40.21 1.00 43.34 1.00 39.06 1.00 42.15 1.00 39.39 1.00 41.03 1.00 39.39 1.00 42.86 1.00 42.30 1.00 42.30 1.00 42.86 1.00 33.26 1.00 30.90 1.00 27.21 1.00 27.34 1.00 30.07 1.00 20.62 1.00 33.39 1.00 35.36 1.00 34.34 1.00 30.53 1.00 36.64 1.00 47.37 1.00 52.82 1.00 56.09 1.00 54.95 1.00 49.51 1.00 49.51 1.00 49.51 1.00 49.51 1.00 49.51 1.00 47.37 1.00 52.82 1.00 56.09 1.00 54.95 1.00 67.56 1.00 71.37 1.00 58.81 1.00 72.21 1.00 74.67 1.00 71.55 1.00 71.37 1.00 58.81 1.00 58.38 1.00 58.38 1.00 58.38 1.00 58.38	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4291 NE ARG 4293 CZ ARG 4294 NH1 ARG 4297 NH2 ARG 4300 C ARG 4301 O ARG 4302 N ALA 4305 CB ALA 4305 C ALA 4306 C ALA 4307 O ALA 4308 N SER 4310 CA SER 4311 CB SER 4312 OG SER 4314 C SER 4315 O SER 4316 N CYS 4318 CA CYS 4318 CA CYS 4321 CB CYS 4322 CG CYS 4321 CB CYS 4321 CB CYS 4322 CO CYS 4321 CB CYS 4323 C1 NAG 4325 C2 NAG 4327 N2 NAG 4327 N2 NAG 4329 C7 NAG 4327 N2 NAG 4329 C7 NAG 4330 C7 NAG 4331 C8 NAG 4335 C3 NAG 4345 O5 NAG 4346 C6 NAG 4347 C7 NAG 4357 C7 NAG 4358 C7 NAG 4359 C8 NAG 4369 O4 NAG 4370 C5 NAG 4370 C7 NAG	457 457 25.542 457 24.070 457 23.463 457 26.383 457 458 27.208 458 28.102 458 28.141 458 29.513 458 29.513 458 29.513 458 29.711 21A 59.799 21A 60.283 21A 59.237 21A 58.645 21A 58.912 21A 57.580 21A 61.497 21A 61.973 21A 62.599 21A 63.672 21A 63.673 21A 63.672 21A 63.673 21A	73.165 25.322 72.528 27.376 71.878 28.417 73.558 27.226 73.993 28.294 75.532 28.384 76.124 29.390 73.447 28.026 71.628 28.098 6.600 61.811 6.883 60.413 6.588 59.464 7.564 58.793 8.758 58.951 7.139 57.803 6.018 60.112 6.353 61.166 5.328 60.986 6.111 62.591 6.253 61.166 5.328 60.986 6.111 62.591 6.921 62.754 6.596 63.618 7.832 63.218 15.309 72.728 15.273 73.299 14.505 72.479 13.529 73.008 13.190 74.171 12.782 72.101 16.691 73.372 16.699 74.013 17.643 74.096 18.985 73.872 17.517 73.555 16.138 73.560 18.324 74.357 17.942 75.718 19.770 74.983 21.174 74.572	1.00 86.96 1.00 77.80 1.00 77.91 1.00 77.15 1.00 75.70 1.00 77.12 1.00 81.46 1.00 88.76 1.00 95.43 1.00 98.69 1.00100.00 1.00 97.6 1.00 74.89 1.00 74.89 1.00 74.89 1.00 74.89 1.00 74.89 1.00 74.89 1.00 74.73 1.00 73.98 1.00 72.39 1.00 73.46 1.00 73.46 1.00 73.46 1.00 73.46 1.00 73.36 1.00 73.36 1.00 73.36 1.00 73.36 1.00 73.36 1.00 73.36 1.00 69.91 1.00 73.36 1.00 69.91 1.00 63.96 1.00 63.96 1.00 68.52 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.83 1.00 70.82 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83 1.00 70.83	AAAA AAAA AAAAA AAAAA AAAAA AAAAA AAAAA AAAA								

• . • •	335R 336R V) 338N	
	OT 312D 312D 53333 T 4V (344'	•
	Face 3  310T 309K 312D 335R 9M (3165) 3135 318Q 315T 336R 334V (344V) 346Q 343E 338N	
	302C) 31	
	Cleff 2   Fc 264E 305E 305B 282I 300K 319M 282I 300K 318Q 298C (322G)321Q 347F 5G) 346Q	
	12   S 262D 2633 2776E (280) (280)	
	Face 2 259E 2618 256L 26 256F 275Q 276F (274M) 272E 27 0R 270D	Figure 2
	Cleff 1 5) 26E 2551 242E 241 79W 24	
	, (6G) 5F 28Y (27G 54Y 53E 2F 3Y 80K	
	Face 1 10R 8E 32L 30H 58F 56L 8 90F (88V) 8 112R 85	
	Face 1   Cleff   Face 2   Cleff   (12D)   10R   259E   261S   262D   256L   263S   264E   261S   261	







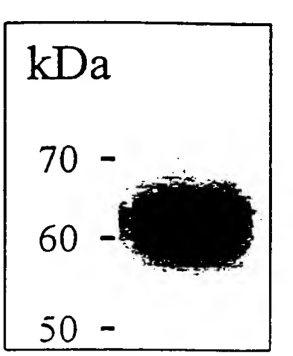


Figure 3

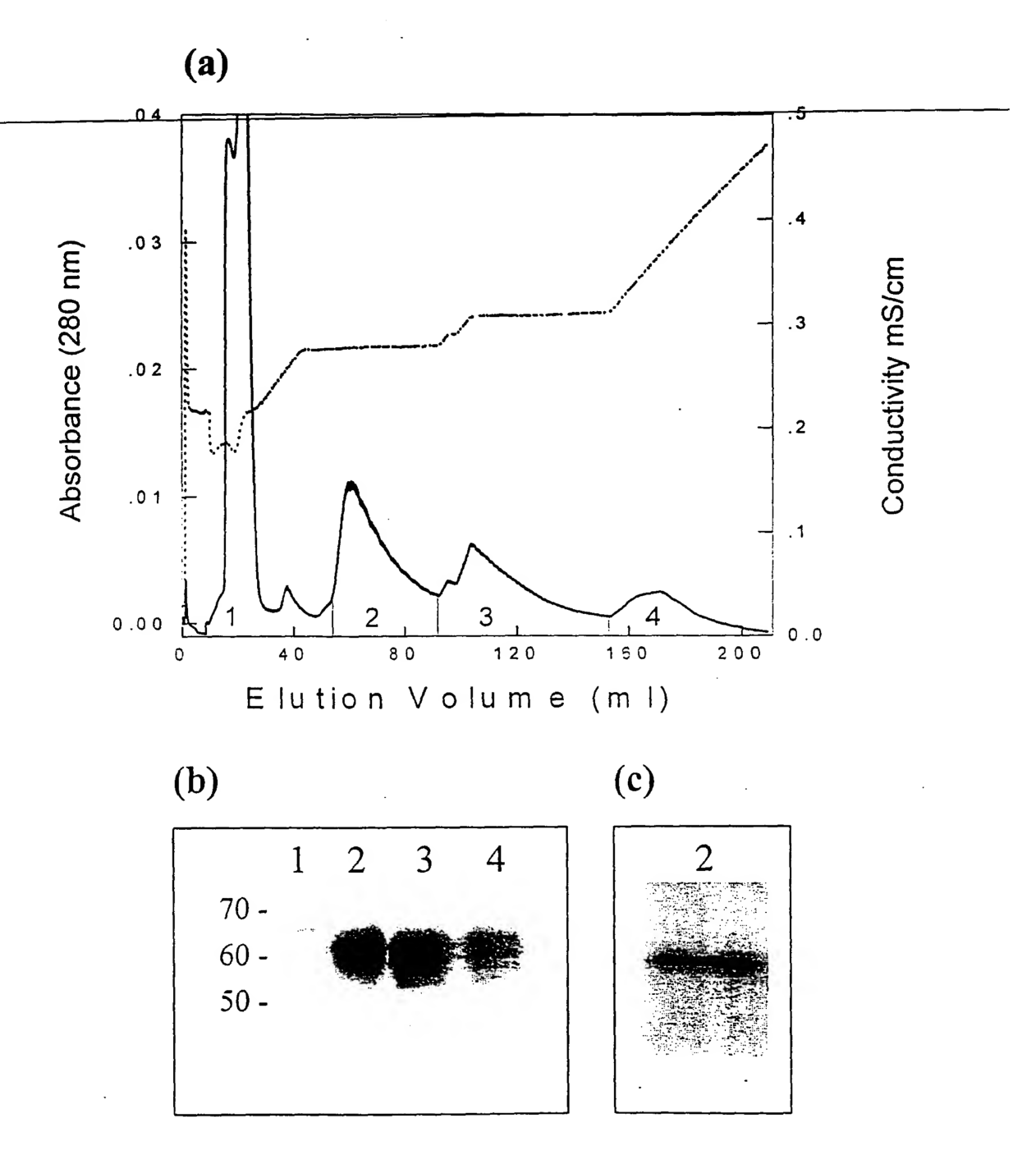
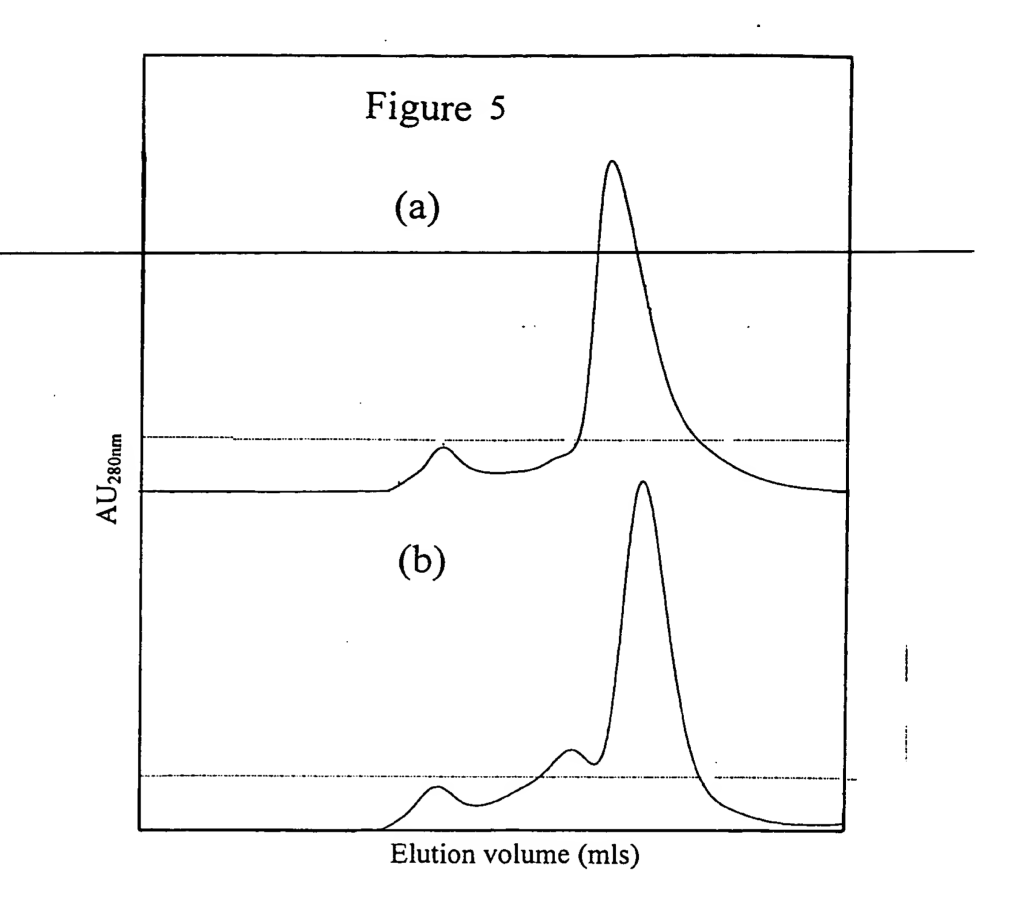
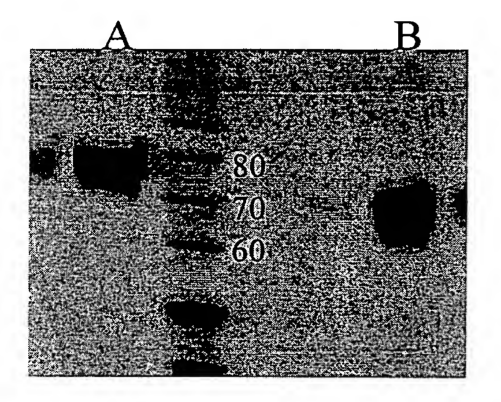


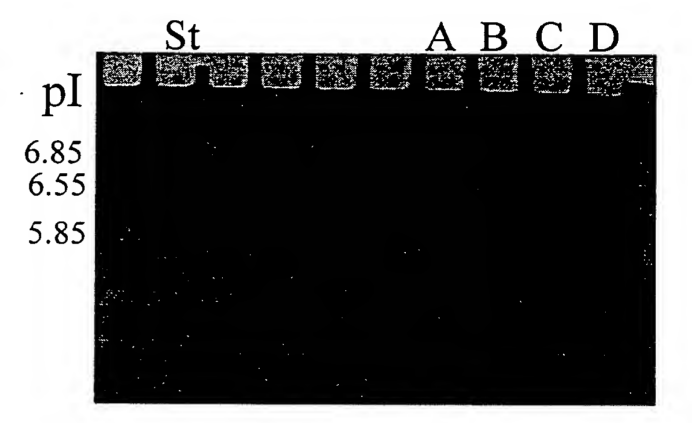
Figure 4



## (a) SDS PAGE



## (b) IEF pH3-7



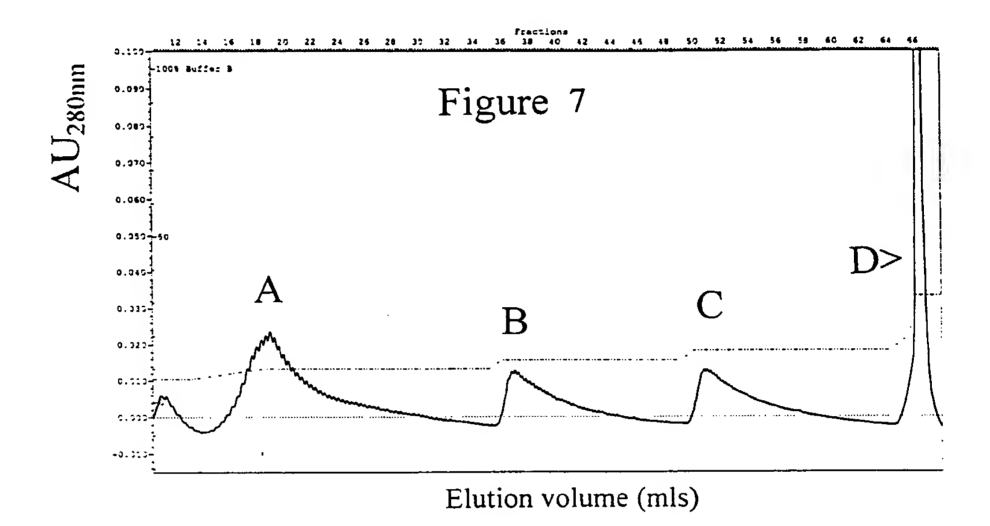
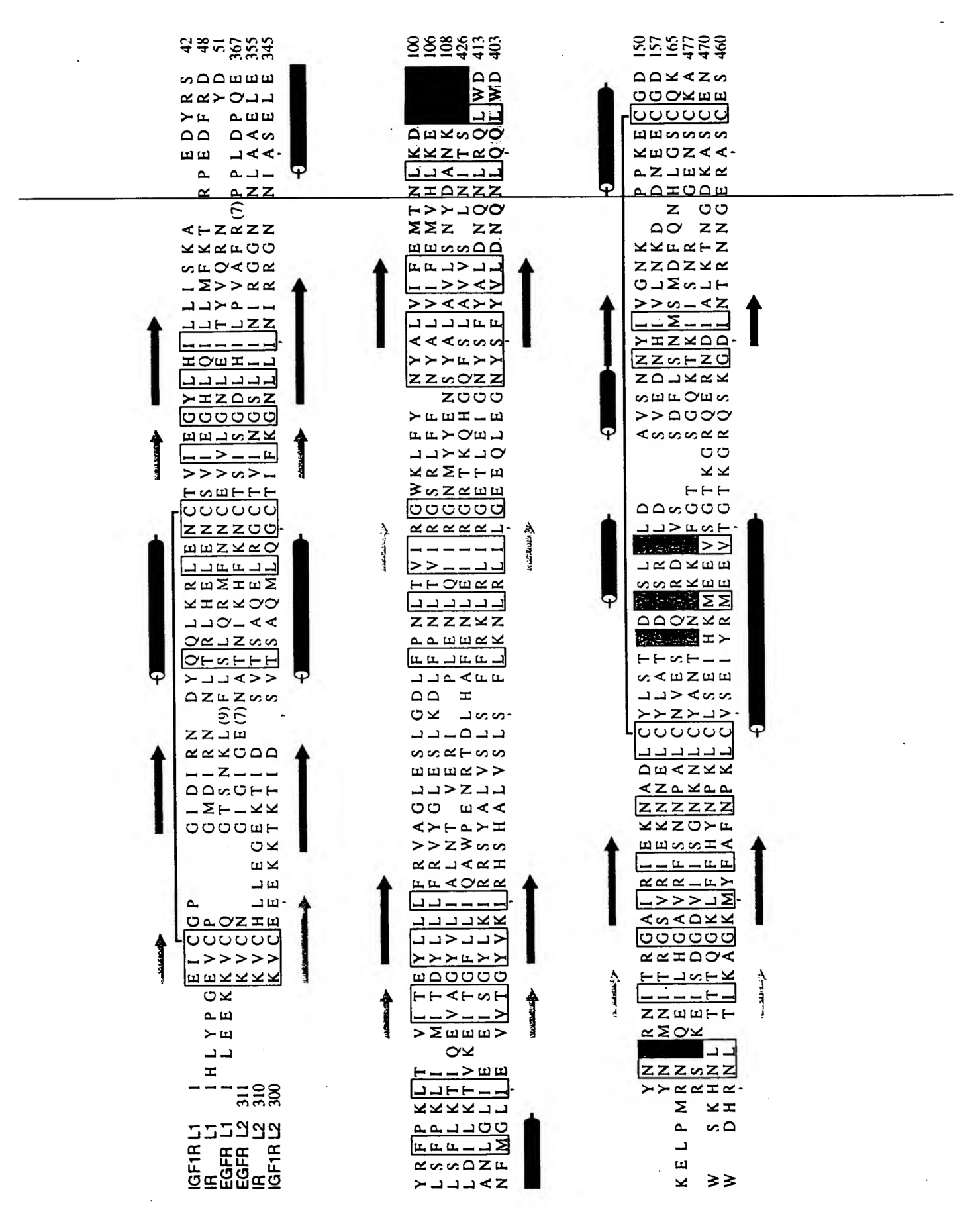
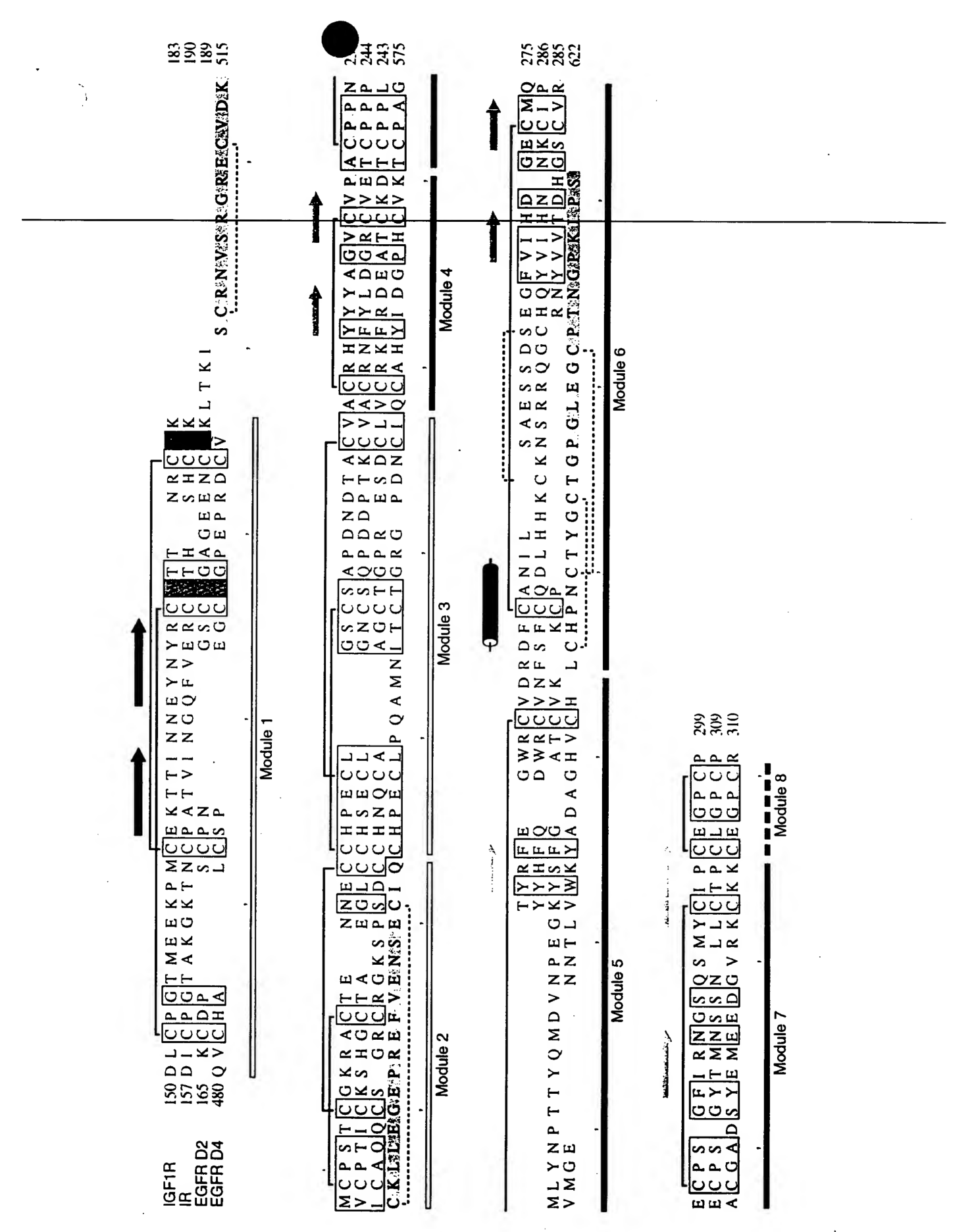




Figure 8





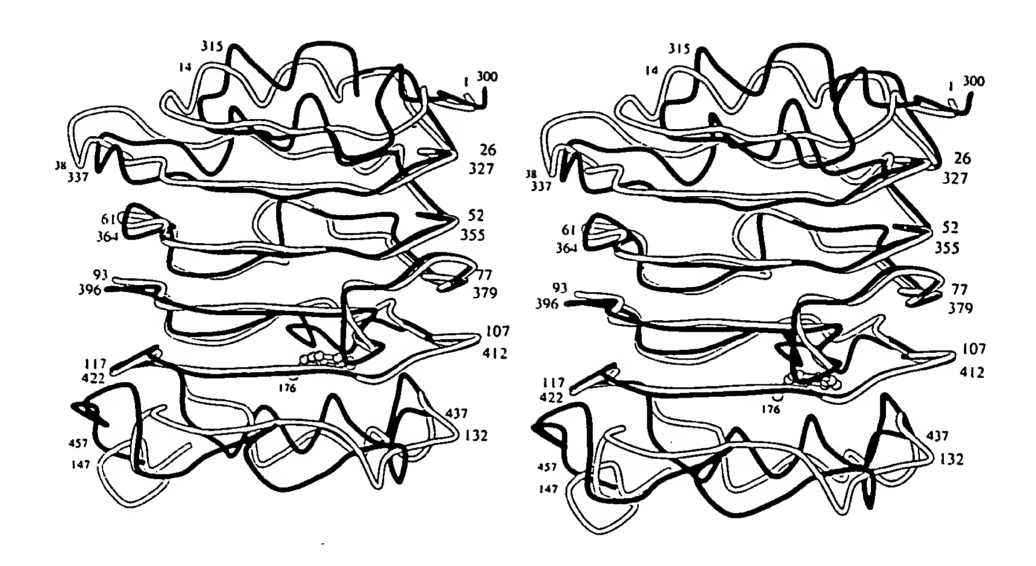
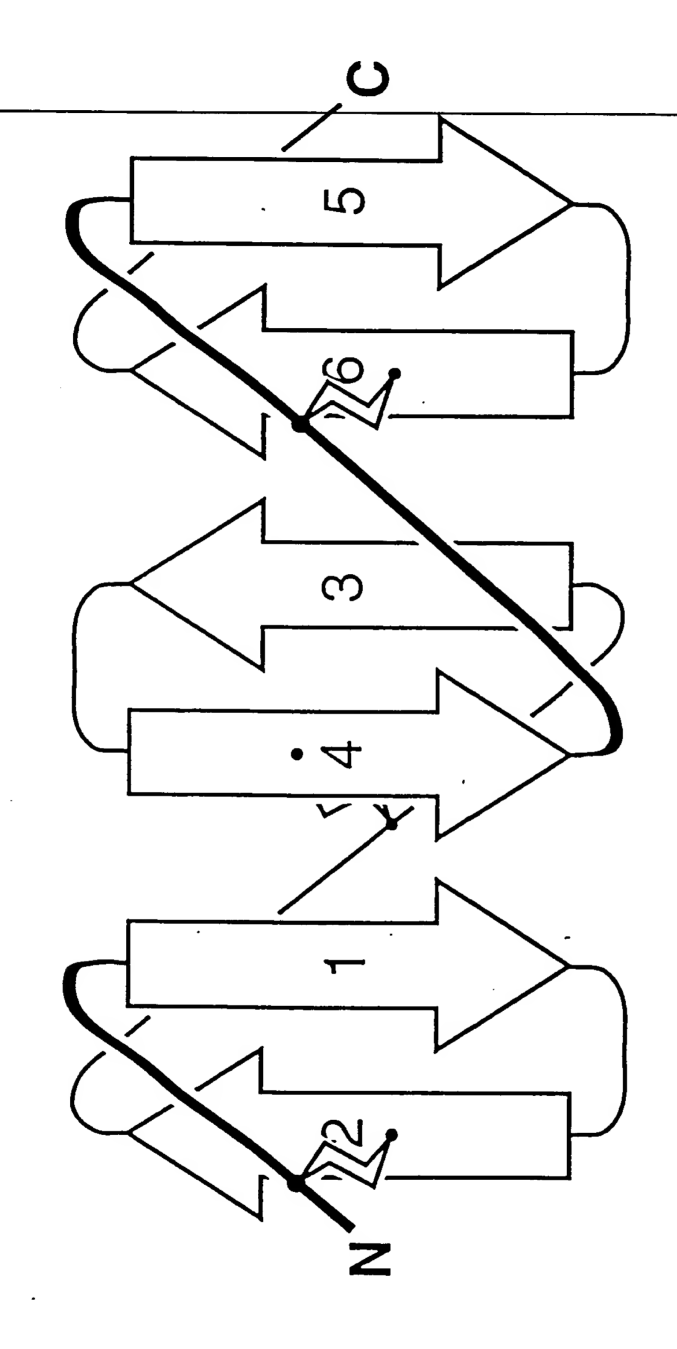
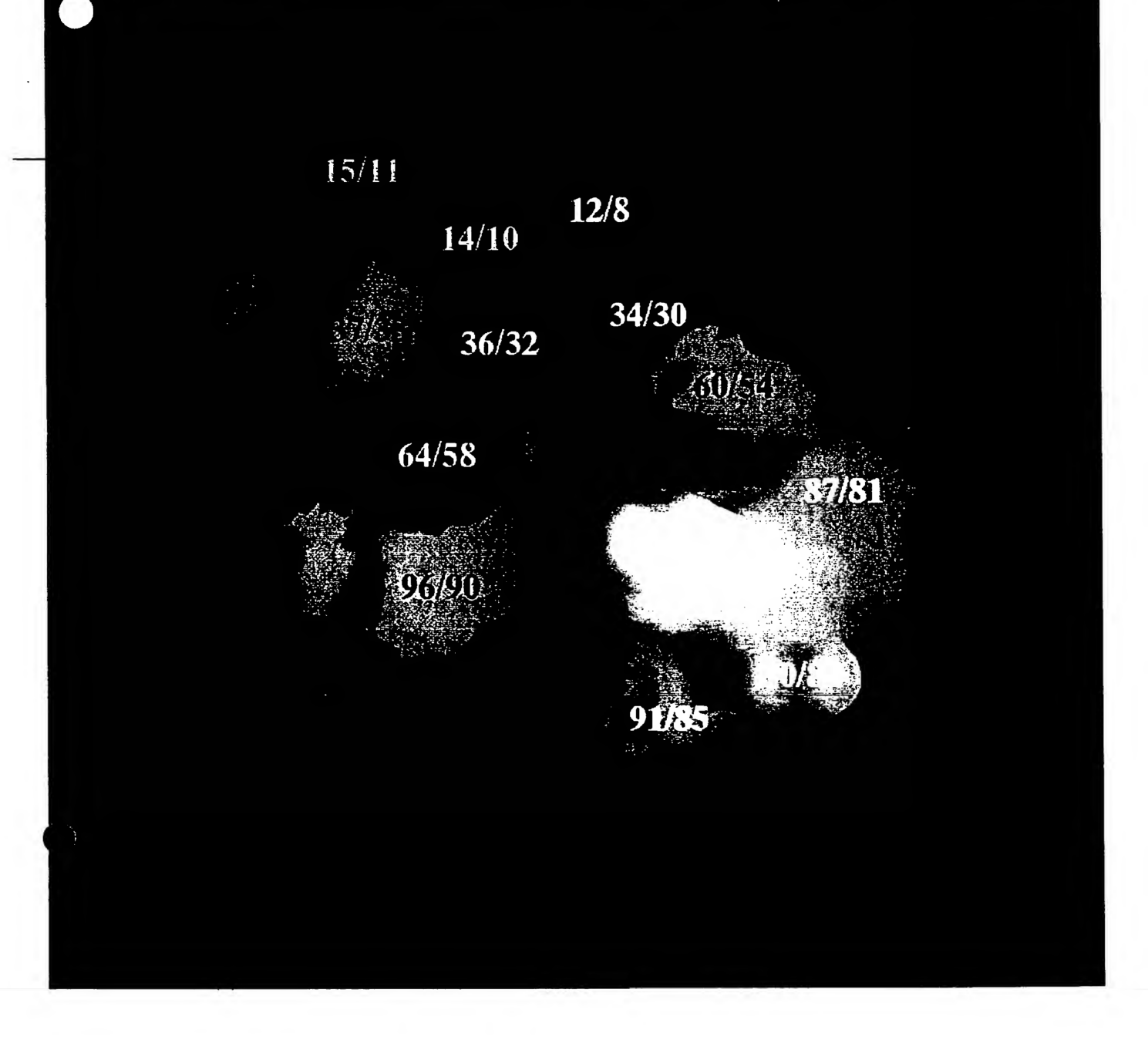


Figure 10





## Figure 13: Sequence Alignment of hIGF-1R, hIR and hIRR ectodomains.

Derived by use of the PileUp program in the software package of the Genetics Computer Group, 575 Science Drive, Madison, Wisconsin, USA.

Symbol Comparison table: GenRunData:PileUpPep.Cmp CompCheCk: 1254

GapWeight: 3.0
<del>GapLengthWeight: 0.1</del>

Name:	Higflr Hir Hirr	Len: Len: Len:	972 <b>C</b> he <b>C</b>	k: 2986 We	ight: 1.00 ight: 1.00 ight: 1.00	
					19110. 1.00	
				•		
11: ~£1 ~	* ET <b>~</b> CD	CIDIDADVOO	*	DOWN HITT T TO	v a repure ou	43
Hir	EICGP		LHELE <u>NCS</u> VI			
	MNVC.P	· —				
				-	•	
Higf1r	סבטעו שעודשב	VIII EDWACI	ECI CDI EDNI	MUTDOWN EV	NYALVIFEMT	0.2
Hir				<del></del>	NYALVIFEMV	
	SFPRLTQVTD			<del></del>		
Higf1r	NUKDIGLYNL	RNTTRGATET	EKNADL <b>C</b> YLS	TVDWSLTLDA	VSNNYIVGNK	143
Hir					VEDNYIVLNK	
Hirr	HLRDVALPAL	GAVLRGAVRV	EKNQEL <i>C</i> HLS	TIDWGLLQPA	PGANHIVGNK	145
	* *		*	* *	* *	
Hiaf1r	PPK.ECGDLC	PGTMEEKPM.				191
-	DDNEE CGDI C				QKVCPTICKS	
Hirr	LG.EECADVC	PGVLGAAGEP	<b>C</b> AKTTFSGHT	DYR <b>C</b> WTSSHC	QRVCPCPHG.	193
	* **	* *	* *	*	*	
Higf1r	RACTENNECC	HPECLGSCSA		RHYYYAGV <i>C</i> V	PACPPNTYRF	241
Hir					ETCPPPYYHF	248
Hirr	MACTARGECC	HTECLGGCSQ	PEDPRACVAC	RHLYFQGA <b>C</b> L	WA CPPGTYQY	243
	*	* *	*	* *		
Higf1r	EGWR <i>C</i> VDRDF	CANILSAES.	SDSEGFV	IHDGE CMQE C	PSGFIRNGSQ	287
Hir	QDWR <i>C</i> V <u>NFS</u> F	_			PSGYTM <u>NSS</u> N	
Hirr	ESWR <i>C</i> VTAER	<b>C</b> ASLHSVPG.	RASTFG	IHQGS <b>C</b> LAQ <b>C</b>	PSGFTR <u>NSS</u> .	287
	* *	* *		*		
Higf1r	SMYCIPCEGP	<b>C</b> PKV <b>C</b> EEEKK	TKTIDSVTSA	QMLQGCTIFK	GNLLINIRRG	337
Hir				<del>-</del>	<u>GS</u> LIINIRGG	
Hirr	SIF <b>C</b> HK <b>C</b> EGL	CPKECKVG	TKTIDSIQAA	QDLVG <b>C</b> THVE	GSLILNLRQG	335
		•				
Higf1r	NNIASELENF	MGLIEVVTGY	VKIRHSHALV	SLSFLKNLRL	ILGEEQLEG <u>N</u>	
Hir			LKIRRSYALV			397
Hirr	YNLEPQLQHS	LGLVETTTGF.	LKIKHSFALV	SLGFFKNLKL	IRGDAMVDG <u>N</u>	385
				*		
Higf1r	<u>YS</u> FYVLDNQN	_				437
Hir	<del></del>	_ <del>-</del>	NLTITQGKLF			447 435
Hirr	TTP1 APDIMÓIN	ロググアG2MAWW	GLTIPVGKIY	FAFNPRL <b>C</b> LE	ULIVIEEAIG	400
				of 1-462 fr	_	
Higflr	<del></del>	NTRNNGERAS			II TWHRYRPPI	
Hir Hirr		ALKTNGDQAS NPRTNGDRAA	- <del>-</del>		LL RWEPYWPPI LL RWERYEPLI	
79 T T T	TWOKÖMKAET	*** IVIIAGDIVAN	exitt nur	A TATURDICE		

Higflr Hir Hirr	RDLLGFMLFY	KEAPFK <u>NVT</u> E KEAPY <u>QNVT</u> E KESPF <u>QNAT</u> E	FDGQDA CGSN	SWTVVDIDPP	PNKDV LRSNDPKSQN LSRTQ	532 547 530
Higflr Hir Hirr	HPGWLMRGLK	PWTQYAVYVK PWTQYAIFVK PWTQYAVFVR	TL.VTFSDER	RTYGAKSDII	YVQTDATNPS	582 596 580
Higf1r Hir Hirr	VPLDPISVS <u>N</u>	SSSQLIVKWN SSSQIILKWK SSSHLLVRWK	PPSDPNG <u>NIT</u>	HYLVFWERQA	EDSELFELDY	632 646 630
Higf1r Hir Hirr	<b>C</b> LKGLKLPSR	KYADGTIDIE TWS.PPFESE N.NDPRFDGE	EVTENPKTEV DSQKH <u>NOS</u> E. DGDPEAEME.	YEDSAGE <i>CC</i> S	* CPKTEAE CPKTDSQ CQHPPPGQVL	678 691 673
Higflr Hir Hirr	KQAEKEEAEY ILKELEESSF PPLEAQEASF			RRSLGDVGNV	TVAVPTV	728 738 722
Higflr Hir Hirr	AAFP <u>NTS</u> STS	DPEELETEYP VPTSPEEHRP DFEIQEDKVP	FEKVVNKE	SLVISGLRHF	TGYRIELQA $c$	786
Higf1r Hir Hirr	NQDTPEER CS	VAAYVSARTM	PEAKADDIVG	PVTHEIFENN	SIFLKWPEPE VVHLMWQEPK SVLLRWLEPP	836
Higflr Hir Hirr	EPNGLIVLYE	VSYRRYGDEE	LHL <b>C</b> VSRKHF	$\mathtt{ALERG} \textbf{\textit{C}} \mathtt{RLRG}$	LNPG <u>NYT</u> ARI LSPG <u>NYS</u> VRI LPPG <u>NYS</u> ARV	875 886 864
Higflr Hir Hirr	RATSLAG <u>NGS</u>	WTDPVFFYVQ WTEPTYFYVT WTDSVAFYIL	DYLDVPSNIA	K		906 917 895

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## Figure 14: Sequence Alignment of EGFR, ErbB2, ErbB3 and ErbB4 Ectodomains.

[For alignment on the IGF-1R fragment see Fig. 9]

Derived by use of the PileUp program in the software package of the Genetics Computer Group, 575 Science Drive, Madison, Wisconsin, USA.

Symbol comparison table: GenRunData: Pileuppep.Cmp CompCheck: 1254

GapWeight: 3.000 GapLengthWeight: 0.100

Name: Name: Name: Name:	: Erb4 : Egfr	Len Len Len Len	: 649 Ch	neck: 790 neck: 2381	Weight: 1.00 Weight: 1.00 Weight: 1.00 Weight: 1.00
Erb3 Erb4 Egfr Erb2	SDSQSVC	AGTENKLSSL QGTSNKLTQL	SDLEQQYRAL GTFEDHFLSI		MGNLEITSIE LGNLEITYVQ
Erb3 Erb4 Egfr Erb2	51 HNADLSFLQW HNRDLSFLRS RNYDLSFLKT TNASLSFLQD	-		ENLRIIRGTK ENLQIIRGNM	100 VYDGKFAIFV LYEDRYALAI YYENSYALAV LFEDNYALAV
Erb3 Erb4 Egfr Erb2	101 MLNYN FLNYR LSNYD LDNGDPLNNT	KDGNFG	LQELGLKNLT LKELPMRNLQ	EILSGGVYIE EILNGGVYVD EILHGAVRFS EILKGGVLIQ	QNKFLCYADT NNPALCNVES
Erb3 Erb4 Egfr Erb2	151 IDWRDIVRDR IHWQDIVRNP IQWRDIVSSD ILWKDIFHKN	WPSNLTLVST	NGSSGCGRCH NHLGSCQKCD	EVC.KGRCWG KSC.TGRCWG PSCPNGSCWG PMCKGSRCWG	200 PGSEDCQTLT PTENHCQTLT AGEENCQKLT ESSEDCQSLT
Erb3 Erb4 Egfr Erb2	RTVCAEQCDG KIICAQQCSG	RCYGPYVSDC RCRGKSPSDC	CHRECAGGCS CHNQCAAGCT	GPQDTDCFAC GPKDTDCFAC GPRESDCLVC GPKHSDCLAC	MNFNDSGACV RKFRDEATCK
Erb3 Erb4 Egfr Erb2	251 PRCPQPLVYN TQCPQTFVYN DTCPPLMLYN LHCPALVTYN	KLTFQLEPNP PTTFQLEHNF PTTYQMDVNP TDTFESMPNP	NAKYTYGAFC EGKYSFGATC	VASCPHNFVV VKKCPHNFVV VKKCPRNYVV VTACPYNYLS	.DSSSCVRAC TDHGSCVRAC
Erb3 Erb4 Egfr Erb2	301 PPDKMEV.DK PSSKMEV.EE GADSYEM.EE PLHNQEVTAE	NGIKMCKPCT DGVRKCKKCE	GLCPKACEGT DICPKACDGI GPCRKVCNGI KPCARVCYGL	GTGSLMSAQT GIGEFKDSLS	350 VDSSNIDGFV VDSSNIDKFI INATNIKHFK VTSANIQEFA
Erb3 Erb4 Egfr Erb2		ILPVAFRGDS	YNAIEAIDPE FTHTPPLDPQ	KLNVFRTVRE KLNVFRTVRE ELDILKTVKE QLQVFETLEE	ITGFLLIQAW
Erb3 Erb4				MKNLNVTSLG LKQQGITSLQ	

Egfr Erb2				VS.LNITSLG LQGLGISWLG	
Erb3 Erb4 Egfr	NIYITDNSNL	CYHHSLNWTK CYYHTINWTT CYANTINWKK	LF.STINQRI	End L2 doma DIKHNRPRRD VIRDNRKAEN KIISNRGENS	CVA EGKVCDP CTA EGMVCNH CKA TGQVCHA
Erb2		CFVHTVPWDQ	LFRNP.HQAL	LHTANRPEDE	CVG EGLACHQ
Erb3	501 LCSSGCWGP	GPGOCLSCRN	YSRGGVCVTH	CNFLNGEPRE	
Erb4	LCSSDGCWGP	_	FSRGRICIES		FENGSICVEC
Egfr	LCSPEGCWGP	EPRDCVSCRN	VSRGRECVDK	CKLLEGEPRE	FVENSECIQC
Erb2	LCARGHCWGP	GPTQCVNCSQ	FLRGQECVEE	CRVLQGLPRE	YVNARHCLPC
	551				600
Erb3		TATCNGSGSD	TCAQCAHFRD	GPHCVSSCPH	GVLGA.KGP.
Erb4		LLTCHGPGPD		GPNCVEKCPD	GLQGA.NSF.
Egfr	HPECLPQAMN	I.TCTGRGPD	NCIQCAHYID	GPHCVKTCPA	GVMGENNTL.
Erb2	HPECQPQN.G	SVTCFGPEAD	QCVACAHYKD	PPFCVARCPS	GVKPDLSYMP
	601				649
Erb3	IYKYPDVQNE	CRPCHENCTQ			• • • • • • •
Erb4	IFKYADPDRE	_		IYYPWTGHST	LPQHARTPL
Egfr	VWKYADAGHV			PTNGPKIPS.	
Erb2	IWKFPDEEGA	CQPCPINCTH	SCVDLDDKGC	PAEQRASPLT	S

Figure 15. Classification of Cys-rich modules
C2-4 denote modules with the 1-3/2-4 double disulphide bond connections.
C1-2 for the single disulphide bonded modules and
C1-2t for stabilised beta turn.

## First Cys-rich region C2-4 modules

C2-4 invuuies	2				
		1 2 3 4			
Higflr	152	CPSTMEEKPM-CEKTTIMEYNYRCWTTHRC QKM	184	(lst)	
 Hir	159	CPGTAKGETH-CPATVINGOEVERCHTHSHC OKY	191	(1st)	
Hìrr	154	CPGVLGAAGEPCAKTTFSGHTDYRCWTSSHC QRV	137	(1st)	
Egfr	156	CDPSCPNG-SCWGAG-EENC QKLTKII	190	(1st)	
hĒrb2	174	CSPHCKGS-RCWGES-SEDC QSLTRTV	198	(1st)	
hErb3	167	CHEVCKGRCWGPG-SEDC QTLTKTI	190	(1st)	
hErb4	157	CHKSCTGRCWGPT-ENHC QTLTRTV	190	(1st)	
Higflr	185	CPSTCGK-RACTENNEC	200	(2nd)	
Hìr	192	CPTICKS-HGCTAEGLC	207	(2nd)	
Hirr	138	CPCPHGMACTARGEC	202	(2nd)	
Egfr		CAQQCSGRCRGKS-PSDC	207	(2nd)	
hErb2		CAGGCARCKGPL-PTDC	214	(2nd)	
hErb3		CAPQCNGHCFGPN-PNQC	207	(2nd)	
hErb4		CAEQCDGRCYGPY-VSDC	207	(2nd)	
POISH	191	CWEGODGWC1GF1-43DG	2.57	(2114)	
Higflr	201	CHPECLGSCSAPDNDTAC VA	220	(3rd)	
Hir		CHSECLGNCSQPDDPTKC VA	227	(3rd)	
		CHTECLGGCSQPEDPRAC VA		(3rd)	•
Hirr 		~	222		
Egir		CHIQCAAGCTGPR-ESEC LV	226	(3rd)	
Erb2		CHEQCAAGCTGPK-HSDC LA	233	(3rd)	
hErb3		CHIECAGGCSGPQ-DTDC FA	226	(3rd)	
hErb:	208	CHRECAGGCSGPK-DTCC FA	226	(3rd)	
C1 2 modulos		·			
C1-2 modules			222		
_		CRHYYYAGVC VPA	233	(4th)	
		CRNFYLDGRC VET	240	(4th)	
		CRHLYFQGAC LWA	235	(4th)	
-		CRHFRDEATC MDT	239	(4th)	
		ClhfnHSGIC ELH	246	(45h)	
hErb3	227	CAHFUDSGAC VPR	239	(4th)	
hBrb4	227	CHRENDSGAC VTQ	239	(ath)	
Higflr	234	CSFNTYREEGWRC VDRDF	251	(5th)	
Hir	141	CREETYHEQDWRC VNESE	259	(5th)	
Hirr	235	CFFSTYQYESWRC VTAER	253	(5th)	
Egfr	249	CFFLMLYHETTYQHDVHPEGKYSFGATC VKK		(5th)	
		CPALVTYHTDTFESHPNPEGRYTFGASC VTA	277	(5th)	
hErc3	240	CPQPLVYNKLTFQLEPNPHTKYQYGGVC VAS		(5th)	
hErb4	240	CPQTEVINETTEQLEHNENAKYTIGAEC VKK	270	(5th)	
114 m 51 m	252	CONTRACTOR CONTRACTOR NOT	276	(6+5)	
_		CAMILSAESSDSEGFVIHD.GEC MQE CQD.LHHKCKNSRRQGCHQYVIHN.NKC IPS		(6th) (6th)	
Hir		CAS.LHSVPGRASTFGIHQ.GSC LAQ		(6th)	
		CPRHYVVTDHGSC VRA	286	(Sth)	
<u>-</u>		CFYNYLSTDVGSC TLV	293	(6th)	
		CEHMEVV. DOTSC VRA	285	(6th)	
hErb4		CEHNEVY. DSSSC VRA	285	(6th)	
				_	
_		CPSG. FIRNGSQ-SNYC IP	293	(7th)	
		CPSG. YTHINSSNLLC TP	303	(7th)	
Hirr		CPSG. FTRIISSSIFC HK	293	(7th)	
Egf:		CGADSYEME + EDGVRKC KK	304	(7th)	
hErbl		C91HNQEVTAEDGTQRC EK	312	(7th)	
hErb3		CRECKIEVDKH-GLKIC EP	303	(7th)	
hErc i	548	CESSMIEVEEN-GIMIC KP	303	(7th)	

C1-2t module			
Higf1r	294 CEGPC	298	(8th)
Hir	304 CLGPC	308	(8th)
Hirr	294 CEGLC	298	(8th)
	305 CEGPC		
hEgfr		. 309	(8th)
hErb2	313 CSKPC	317	(8th)
hErb3	304 CGGLC	308	(8th)
hErb4	304 CTDIC	308	(8th) -
Second Cys-rich	region.		
C2-4 modules			
hEgfr	482 CHALCSPEGCWGPEPRDCVS	501	(1st)
hErb2	490 CHQLCARGHCWGPGPTQCVN	509	(1st)
hErb3	481 CDPLCSSGGCWGPGPGQCLS	500	(1st)
hErb4	481 CNHLCSSDGCWGPGPDQCLS	500	(1st)
Egfr	534 CHPECLPQAM-NITCTGRGPDNC IQ	557	(4th)
hErb2	542 CHPECQPQNG-SVTCFGPEADQC VA	565	(4th)
hErb3	533 CHPECQPMEG-TATCNGSGSDTC AQ	556	(4th)
hErb4	533 CDPQCEKMEDGLLTCHGPGPDNC TK	557	(4th)
honen	596 CHPNCTYGCTGPGLEGC PTNGPKIPS/	621	(7th)
hEgfr hErb2	605 CPINCTHSCVDLDDKGC PAEQRAQRASPLTS/	632	(7th)
hErb3	594 CHENCTOGCKGPELQDC LGQT/	614	(7th)
hErb4	595 CHPNCTQGCNGPTSHDC IYYPWTGHSTLPQHAI	RTPL 630	(7th)
C1-2 modules			
hEgfr	502 CRNVSRGREC VDK	514	(2nd)
hErb2	510 CSQFLRGQEC VEE	522	(2nd)
	501 CRNYSRGGVC VTH	513	(2nd)
hErb4	501 CRRFSRGRIC IES	513	(2nd)
hEgfr	515 CKLLEGEPREFVENSEC IQ	533	(3rd)
hErb2	523 CRVLQGLPREYVNARHC LP	541	(3rd)
hErb3	514 CNFLNGEPREFAHEAEC FS	532	(3rd)
hErb4	514 CNLYDGEFREFENGSIC VE	532	(3rd)
		570	(544)
_	558 CAHYIDGPHC VKT	570	(5th)
hErb2	566 CAHYKDPPFC V-A	578	(5th)
hErb3	557 CAHFRDGPHC V-S	569	(5th)
hErb4	558 CSHFKDGPNC VEK	570	(5th)
hEgfr	571 CPAGVMGENNTL-VWKYADAGHVC HL	595	(6th)
hErb2	579 CPSGVKPDLSYMPIWKFPDEEGAC QP	604	(6th)
hErb3	570 CPHGVLGAKGPIYKYPDVQNEC RP	593	(6th)
hErb4	571 CPDGLQGANSFIFKYADPDREC HP	594	(6th)
See Pattern is:		C1 25	
IR family: EGFR family:1			
_	C2-4, $C2-4$ , $C2-4$ , $C1-2$	., CI-26	
•	C2-4, C1-2, C1-2,		
	C2-4		

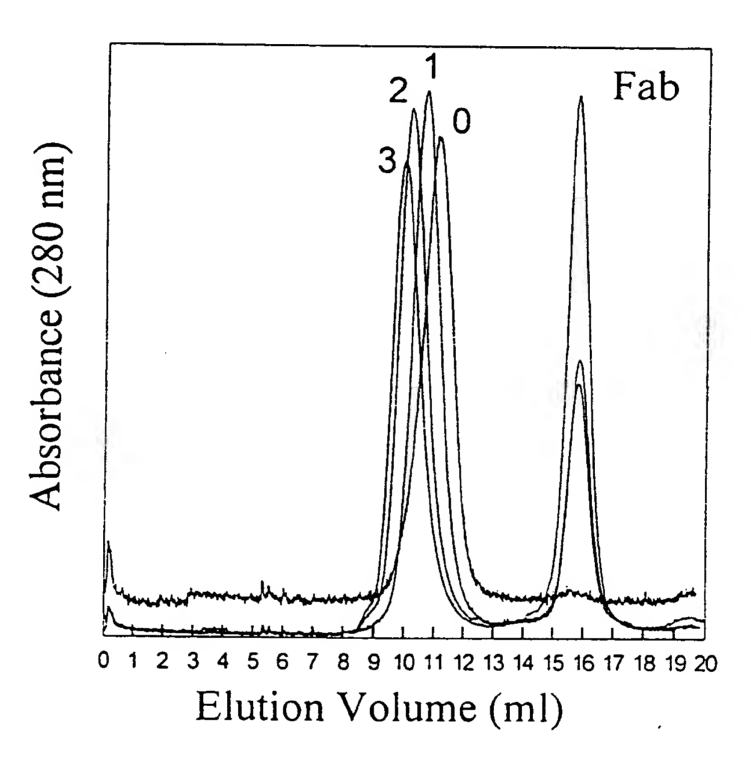
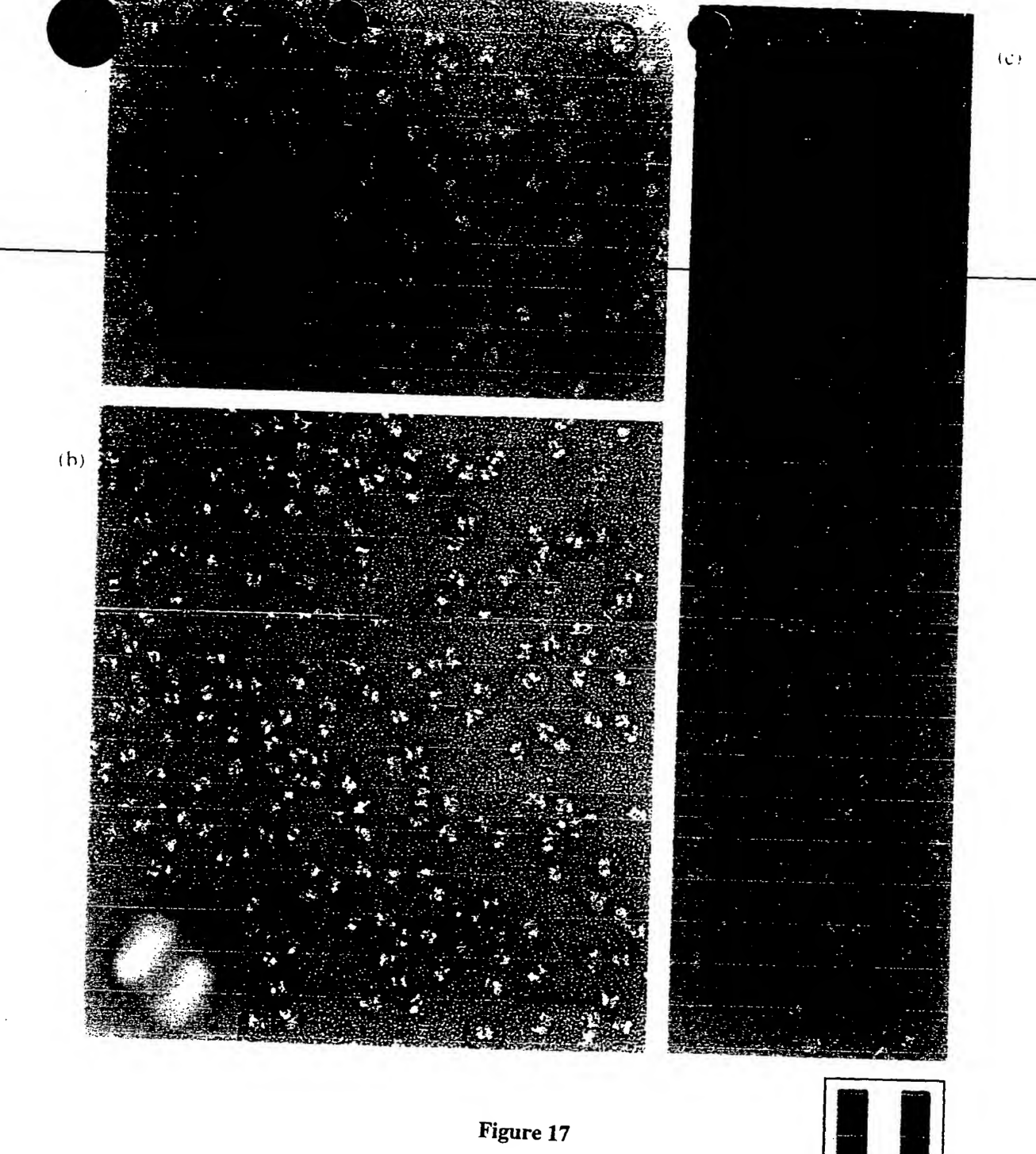
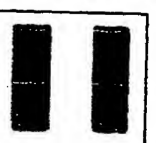


Figure 16





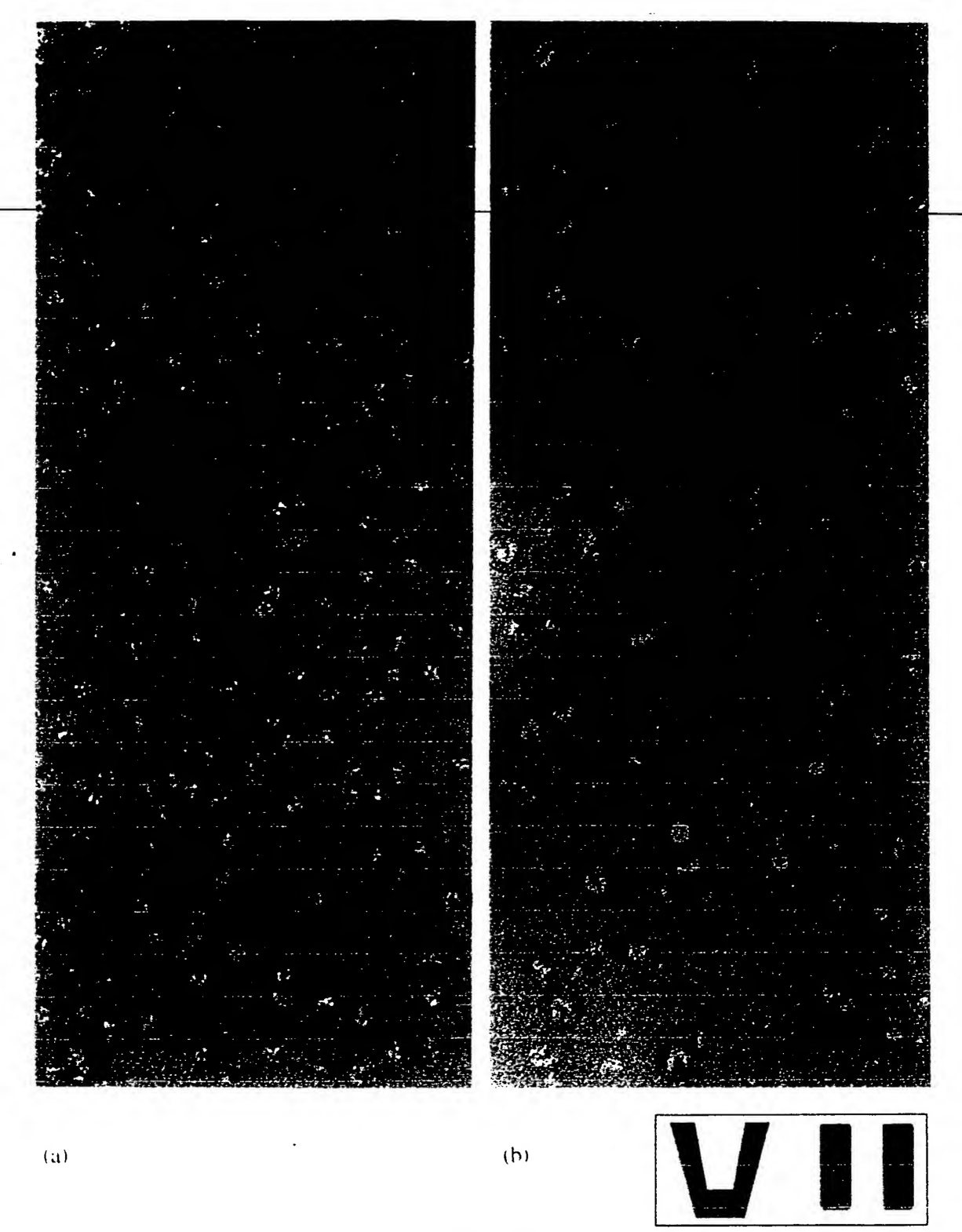


Figure 18

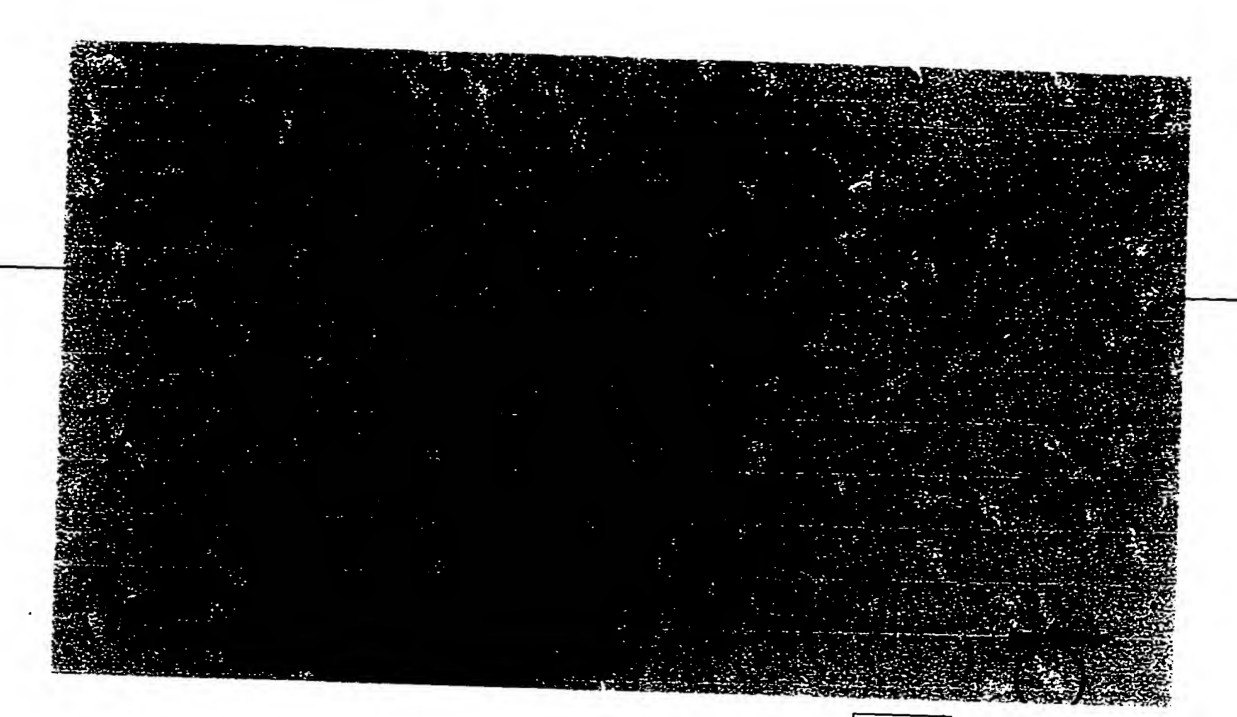
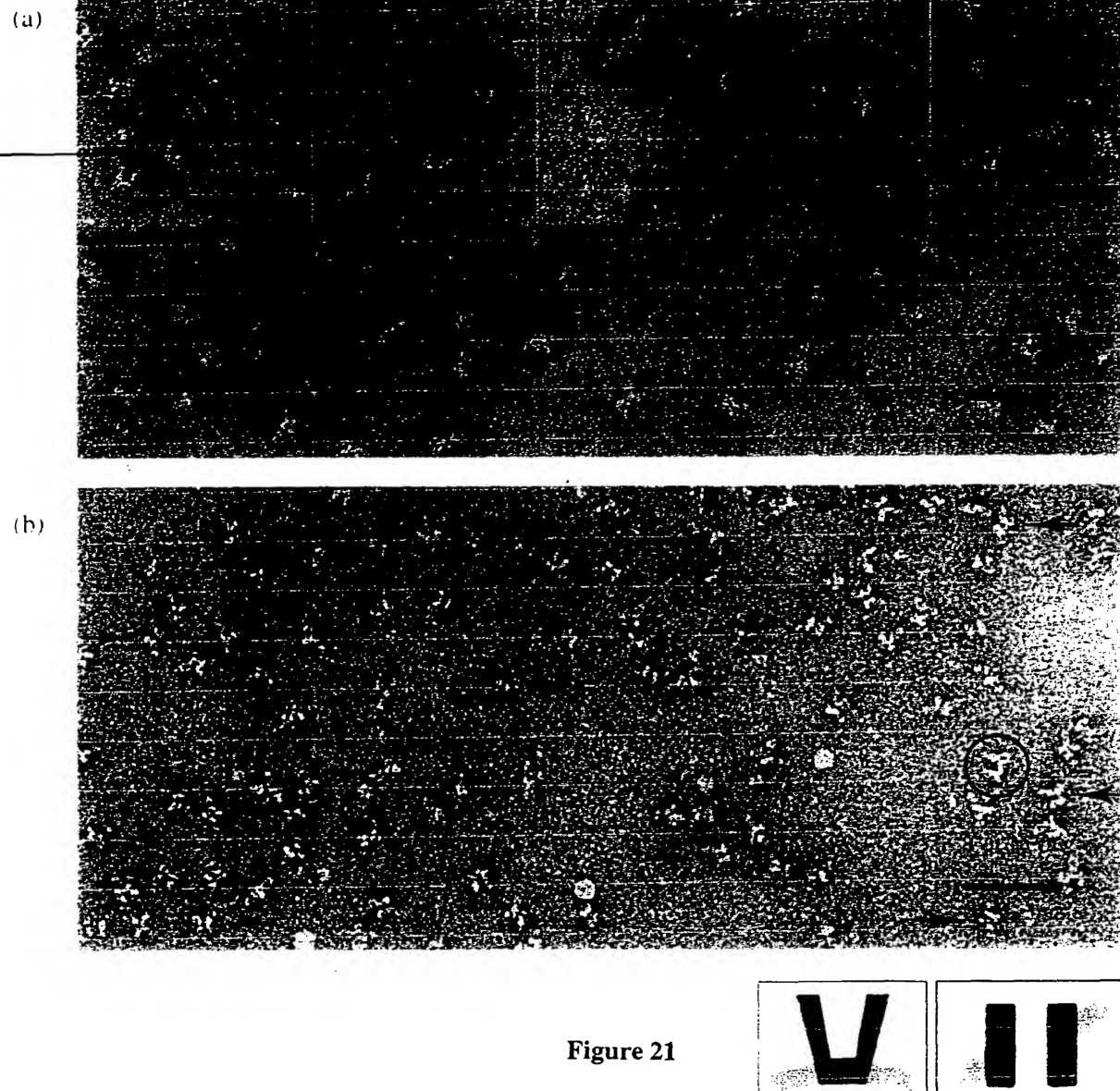


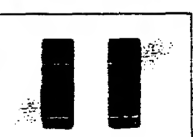
Figure 19



Figure 20







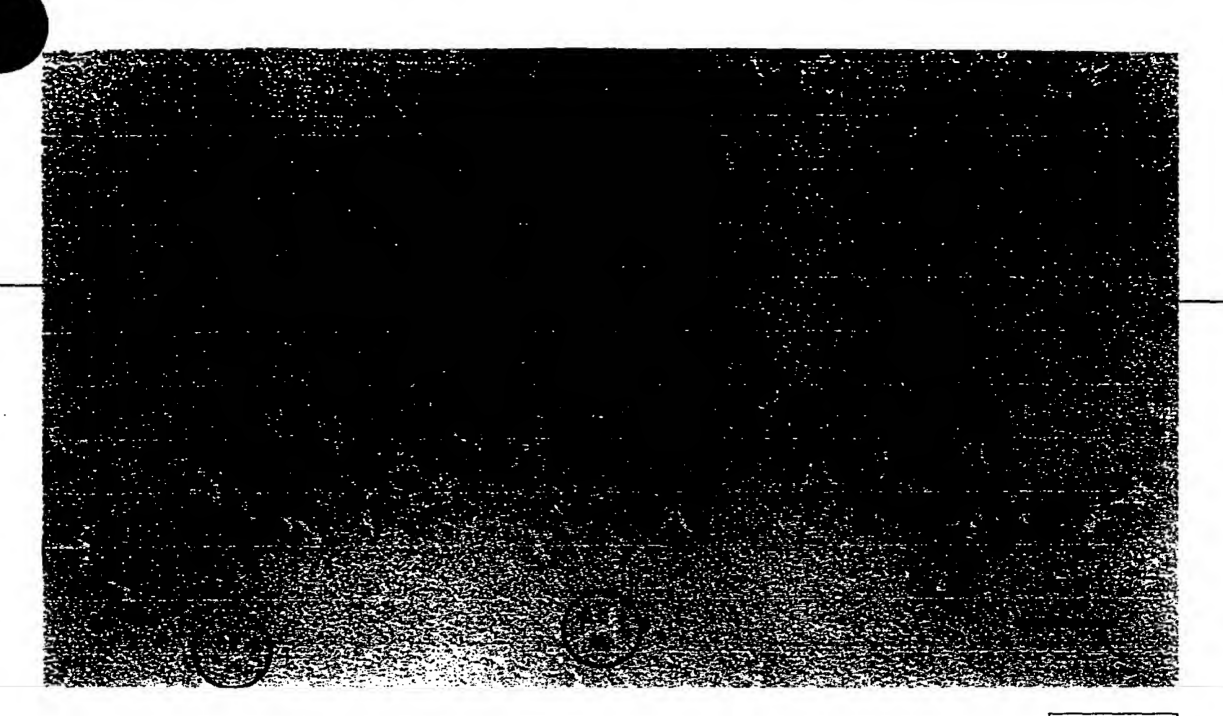


Figure 22



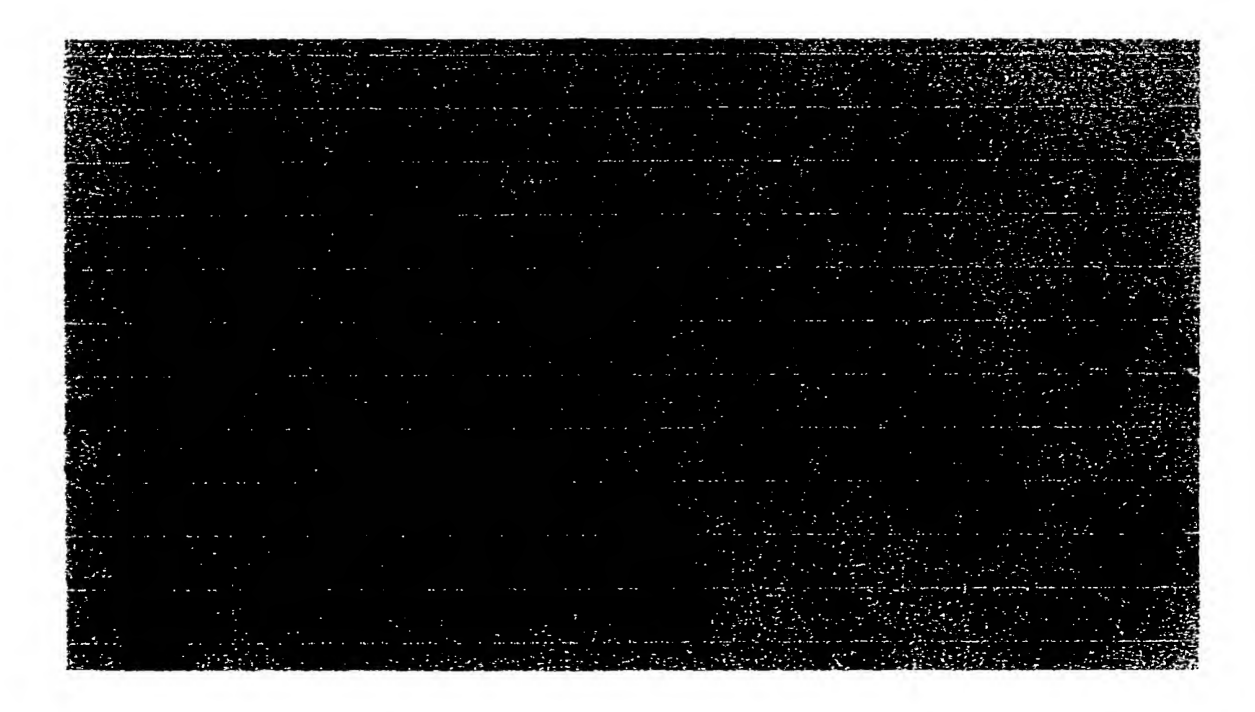


Figure 23

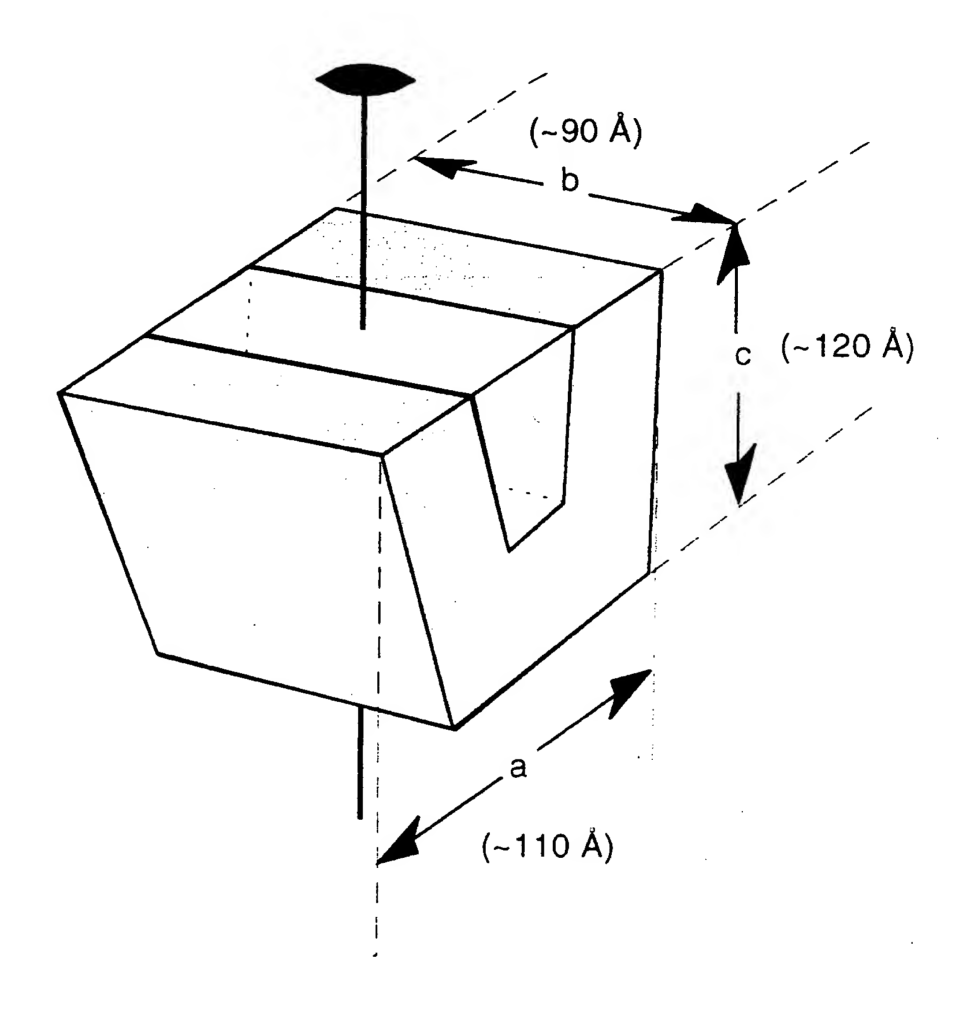


Figure 24

